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## TITLE OF THE INVENTION:

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# NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO M. CATARRHALIS FOR DIAGNOSTICS AND THERAPEUTICS

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### RELATED APPLICATIONS:

This application claims the benefit of U.S. Provisional Application Serial Number 60/128,476, filed April 9, 1999, the entire teachings of which are incorporated herein by reference.

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#### BACKGROUND OF THE INVENTION

The genus Moraxella is a member of the family Neisseriaceae. The 10 species of this genus are separated into 2 subgenera, Moraxella (rods) and Branhamella (cocci). Moraxella are gram-negative, aerobic, oxidase-positive, and usually catalase-postive. (Bovre, K. 1984. Genus II. Moraxella Lwoff 1939, 173 emend. Henriksen and Bovre 1968, 391, 105. Krieg and Holt (editors) In Bergey's Manual of Systematic Bacteriology, 1:296-303.). Moraxella catarrhalis, a member of the subgenera Branhamella, was previously called Branhamella catarrhalis and Neisseria catarrhalis.

Moraxella catarrhalis is frequently isolated from the nasal cavity of humans, and until recently, was considered a nonpathogenic commensal of the upper respiratory tract. Currently it is most important lower respiratory pathogen after S. pneumoniae and H. influenzae (Doren, G., et al, 1986. Diagn. Microbiol. Infect. Dis. 4:191-201.). It is a common cause of otitis media in children, acute bronchitis or pneumonia in adults, and sinusitis (Wood, G., et al, 1996. Clin. Infect. Dis. 22:632-636.). Bacteremia, meningitis, skeletal infections and endocarditis due to M. catarrhalis are rare, but are observed in immunocompromised individuals (Aebi, C., et al, 1998. Infect. Immun. 66:540-548.). Concern for M. catarrhalis infections of cystic fibrosis (CF) patients is growing. Damage to the respiratory tract by M. catarrhalis could promote invasion by other pathogens such as P. aeruginosa in CF patients. (Deneuville, E., et al, 1995. ACTA Paediatr. 84:1212.). M. catarrhalis is also associated with acute laryngitis. In one study, 50% of patients with acute laryngitis were colonized with M. catarrhalis (Hol, C., et al, 1996. Journal of Infectious Diseases. 174:636-638.), while isolates from healthy adults occur at the rate of 6% -11%. The colonization rates of children can be much higher, with average rates of 30%-35% (Sehgal, SC. et al, 1994. Infection 22:193-196.). In some hospitals, M. catarrhalis accounts for half of all the respiratory infections (Bluesone, C., et al, 1992. Pediatr. Infect. Dis. J. 11:S7-S11.).

Increasing levels of antibiotic resistance have been observed in clinical isolates of *M. catarrhalis* recently. Before 1980, less than 10% of *M. catarrhalis* isolates were ß-lactamase-positive. Currently, most clinical isolates produce ß-lactamase, making them resistant to ß-lactam antibiotics such as penicillin. (Doern, G., *et al*, 1996. Antimicob.

Agents Chemother. 40:2884-2886.). *M. catarrhalis* is intrinsically resistant to a small group of drugs that include vancomycin and trimethoprim (Wallace, RJ. 1990. Am. J. Med. 88:46S-50S), and is becoming increasingly resistant to sulfamethoxazole, oral cephalosporins, and macrolides (Hoppe, HL. 1998. Am. J. Health. Syst. Pharm. 55:1881-97).

Although, *M. catarrhalis* was once considered only as part of the nonpathogenic flora of the upper respiratory tract, it is emerging as an important respiratory pathogen. Currently, it is the third leading cause of lower respiratory tract infections and otitis media. Sequencing and further analysis of this genome will aid in identification of essential genes for development of drug targets, and reduce the health threat this organism poses.

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#### SUMMARY OF THE INVENTION

The present invention fulfills the need for diagnostic tools and therapeutics by providing bacterial-specific compositions and methods for detecting *Moraxella* species including *M. catarrhalis*, as well as compositions and methods useful for treating and preventing *Moraxella* infection, in particular, *M. catarrhalis* infection, in vertebrates including mammals.

The present invention encompasses isolated nucleic acids and polypeptides derived from *M. catarrhalis* that are useful as reagents for diagnosis of bacterial disease, components of effective antibacterial vaccines, and/or as targets for antibacterial drugs including anti-*M. catarrhalis* drugs. They can also be used to detect the presence of *M. catarrhalis* and other *Moraxella* species in a sample; and in screening compounds for the ability to interfere with the *M. catarrhalis* life cycle or to inhibit *M. catarrhalis* infection. They also have use as biocontrol agents for plants.

In one aspect, the invention features compositions of nucleic acids corresponding to entire coding sequences of *M. catarrhalis* proteins (SEQ ID NO: 1 - SEQ ID NO: 1920), including surface or secreted proteins or parts thereof, nucleic acids capable of binding mRNA from *M. catarrhalis* proteins to block protein translation, and methods for producing *M. catarrhalis* proteins or parts thereof using peptide synthesis and recombinant DNA techniques. This invention also features antibodies and nucleic acids useful as probes to

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detect *M. catarrhalis* infection. In addition, vaccine compositions and methods for the protection or treatment of infection by *M. catarrhalis* are within the scope of this invention.

The nucleotide sequences provided in SEQ ID NO: 1 - SEQ ID NO: 1920, a fragment thereof, or a nucleotide sequence at least about 99.5% identical to a sequence contained within SEQ ID NO: 1 - SEQ ID NO: 1920 may be "provided" in a variety of medias to facilitate use thereof. As used herein, "provided" refers to a manufacture, other than an isolated nucleic acid molecule, which contains a nucleotide sequence of the present invention, i.e., the nucleotide sequence provided in SEQ ID NO: 1 - SEQ ID NO: 1920, a fragment thereof, or a nucleotide sequence at least about 99.5% identical to a sequence contained within SEQ ID NO: 1 - SEQ ID NO: 1920. Uses for and methods for providing nucleotide sequences in a variety of media is well known in the art (see e.g., EPO Publication No. EP 0 756 006).

In one application of this embodiment, a nucleotide sequence of the present invention can be recorded on computer readable media. As used herein, "computer readable media" refers to any media which can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage media, and magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. A person skilled in the art can readily appreciate how any of the presently known computer readable media can be used to create a manufacture comprising computer readable media having recorded thereon a nucleotide sequence of the present invention.

As used herein, "recorded" refers to a process for storing information on computer readable media. A person skilled in the art can readily adopt any of the presently known methods for recording information on computer readable media to generate manufactures comprising the nucleotide sequence information of the present invention.

A variety of data storage structures are available to a person skilled in the art for creating a computer readable media having recorded thereon a nucleotide sequence of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the nucleotide sequence information of the

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Washington, D.C. (1997).

present invention on computer readable media. The sequence information can be represented in a word processing text file, formatted in commercially-available software such as WordPerfect and Microsoft Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A person skilled in the art can readily adapt any number of data processor structuring formats (e.g. text file or database) in order to obtain computer readable media having recorded thereon the nucleotide sequence information of the present invention.

By providing the nucleotide sequence of SEQ ID NO: 1 - SEQ ID NO: 1920, a fragment thereof, or a nucleotide sequence at least about 99.5% identical to SEQ ID NO: 1 -10 SEQ ID NO: 1920 in computer readable form, a person skilled in the art can routinely access the coding sequence information for a variety of purposes. Computer software is publicly available which allows a person skilled in the art to access sequence information provided in a computer readable media. Examples of such computer software include programs of the "Staden Package", "DNA Star", "MacVector", GCG "Wisconsin Package" (Genetics Computer Group, Madison, WI) and "NCBI Toolbox" (National Center For Biotechnology Information). Suitable programs are described, for example, in Martin J. Bishop, ed., Guide to Human Genome Computing, 2d Edition, Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, The Internet and the New Biology: Tools for Genomic and Molecular Research, American Society for Microbiology,

Computer algorithms enable the identification of M. catarrhalis open reading frames (ORFs) within SEQ ID NO: 1 - SEQ ID NO: 1920 which contain homology to ORFs or proteins from other organisms. Examples of such similarity-search algorithms include the BLAST [Altschul et al., J. Mol. Biol. 215:403-410 (1990)] and Smith-Waterman [Smith and 25 Waterman (1981) Advances in Applied Mathematics, 2:482-489] search algorithms. Suitable search algorithms are described, for example, in Martin J. Bishop, ed., Guide to Human Genome Computing, 2d Edition, Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, The Internet and the New Biology: Tools for Genomic and Molecular Research, American Society for Microbiology, Washington, D.C. (1997). Such algorithms are utilized on computer systems as exemplified 30

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below. The ORFs so identified represent protein encoding fragments within the *M.* catarrhalis genome and *M. catarrhalis plasmids* and are useful in producing commercially important proteins such as enzymes used in fermentation reactions and in the production of commercially useful metabolites.

The present invention further provides systems, particularly computer-based systems, which contain the sequence information described herein. Such systems are designed to identify commercially important fragments of the *M. catarrhalis* genome and plasmids. As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A person skilled in the art can readily appreciate that any one of the currently available computer-based systems is suitable for use in the present invention. The computer-based systems of the present invention comprise a data storage means having stored therein a nucleotide sequence of the present invention and the necessary hardware means and software means for supporting and implementing a search means. As used herein, "data storage means" refers to memory which can store nucleotide sequence information of the present invention, or a memory access means which can access manufactures having recorded thereon the nucleotide sequence information of the present invention.

As used herein, "search means" refers to one or more programs which are implemented on the computer-based system to compare a target sequence or target structural motif with the sequence information stored within the data storage means. Search means are used to identify fragments or regions of the *M. catarrhalis* genome and plasmids which are similar to, or "match", a particular target sequence or target motif. A variety of known algorithms are known in the art and have been disclosed publicly, and a variety of commercially available software for conducting homology-based similarity searches are available and can be used in the computer-based systems of the present invention. Examples of such software includes, but is not limited to, FASTA (GCG Wisconsin Package), Bic\_SW (Compugen Bioccelerator), BLASTN2, BLASTP2, BLASTX2 (NCBI) and Motifs (GCG).

Suitable software programs are described, for example, in Martin J. Bishop, ed., Guide to

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Human Genome Computing, 2d Edition, Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, The Internet and the New Biology: Tools for Genomic and Molecular Research, American Society for Microbiology, Washington, D.C. (1997). A person skilled in the art can readily recognize that any one of the available algorithms or implementing software packages for conducting homology searches can be adapted for use in the present computer-based systems.

As used herein, a "target sequence" can be any DNA or amino acid sequence of six or more nucleotides or two or more amino acids. A person skilled in the art can readily recognize that the longer a target sequence is, the less likely a target sequence will be present as a random occurrence in the database. The most preferred sequence length of a target sequence is from about 10 to 100 amino acids or from about 30 to 300 nucleotide residues. However, it is well recognized that many genes are longer than 500 amino acids, or 1.5 kb in length, and that commercially important fragments of the *M. catarrhalis* genome and plasmids from *M. catarrhalis*, such as sequence fragments involved in gene expression and protein processing, will often be shorter than 30 nucleotides.

As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a specific functional domain or three-dimensional configuration which is formed upon the folding of the target polypeptide. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzymatic active sites, membrane-spanning regions, and signal sequences. Nucleic acid target motifs include, but are not limited to, promoter sequences, hairpin structures and inducible expression elements (protein binding sequences).

A variety of structural formats for the input and output means can be used to input and output the information in the computer-based systems of the present invention. A preferred format for an output means ranks fragments of the *M. catarrhalis* genome and plasmids possessing varying degrees of homology to the target sequence or target motif. Such presentation provides a person skilled in the art with a ranking of sequences which contain various amounts of the target sequence or target motif and identifies the degree of homology contained in the identified fragment.

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A variety of comparing means can be used to compare a target sequence or target motif with the data storage means to identify sequence fragments of the *M. catarrhalis* genome and plasmids. In the present examples, implementing software which implement the BLASTP2 and bic\_SW algorithms (Altschul et al., J Mol. Biol. 215:403-410 (1990);

- Compugen Biocellerator) was used to identify open reading frames within the *M. catarrhalis* genome and plasmids. A person skilled in the art can readily recognize that any one of the publicly available homology search programs can be used as the search means for the computer-based systems of the present invention. Suitable programs are described, for example, in Martin J. Bishop, ed., *Guide to Human Genome Computing*, 2d Edition,
- Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, *The Internet and the New Biology: Tools for Genomic and Molecular Research*, American Society for Microbiology, Washington, D.C. (1997).

The invention features M. catarrhalis polypeptides, preferably a substantially pure preparation of an M. catarrhalis polypeptide, or a recombinant M. catarrhalis polypeptide. In preferred embodiments: the polypeptide has biological activity; the polypeptide has an amino acid sequence at least about 60%, 70%, 80%, 90%, 95%, 98%, or 99% identical to an amino acid sequence of the invention contained in the Sequence Listing, preferably it has about 65% sequence identity with an amino acid sequence of the invention contained in the Sequence Listing, and most preferably it has about 92% to about 99% sequence identity with an amino acid sequence of the invention contained in the Sequence Listing; the polypeptide has an amino acid sequence essentially the same as an amino acid sequence of the invention contained in the Sequence Listing; the polypeptide is at least about 5, 10, 20, 50, 100, or 150 amino acid residues in length; the polypeptide includes at least about 5, preferably at least about 10, more preferably at least about 20, still more preferably at least about 50, 100, or 150 contiguous amino acid residues of the invention contained in the Sequence Listing. In yet another preferred embodiment, the amino acid sequence which differs in sequence identity by about 7% to about 8% from the M. catarrhalis amino acid sequences of the invention contained in the Sequence Listing is also encompassed by the invention.

In preferred embodiments: the *M. catarrhalis* polypeptide is encoded by a nucleic acid of the invention contained in the Sequence Listing, or by a nucleic acid having at least

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about 60%, 70%, 80%, 90%, 95%, 98%, or 99% sequence identity or % homology with a nucleic acid of the invention contained in the Sequence Listing.

In a preferred embodiment, the subject *M. catarrhalis* polypeptide differs in amino acid sequence at about 1, 2, 3, 5, 10 or more residues from a sequence of the invention contained in the Sequence Listing. The differences, however, are such that the *M. catarrhalis* polypeptide exhibits an *M. catarrhalis* biological activity, e.g., the *M. catarrhalis* polypeptide retains a biological activity of a naturally occurring *M. catarrhalis* enzyme.

In preferred embodiments, the polypeptide includes all or a fragment of an amino acid sequence of the invention contained in the Sequence Listing; fused, in reading frame, to additional amino acid residues, preferably to residues encoded by genomic DNA 5' or 3' to the genomic DNA which encodes a sequence of the invention contained in the Sequence Listing.

In yet other preferred embodiments, the *M. catarrhalis* polypeptide is a recombinant fusion protein having a first *M. catarrhalis* polypeptide portion and a second polypeptide portion, e.g., a second polypeptide portion having an amino acid sequence unrelated to *M. catarrhalis*. The second polypeptide portion can be, e.g., any of glutathione-S-transferase, a DNA binding domain, or a polymerase activating domain. In preferred embodiment the fusion protein can be used in a two-hybrid assay.

Polypeptides of the invention include those which arise as a result of alternative transcription events, alternative RNA splicing events, and alternative translational and postranslational events.

In a preferred embodiment, the encoded *M. catarrhalis* polypeptide differs (e.g., by amino acid substitution, addition or deletion of at least one amino acid residue) in amino acid sequence at about 1, 2, 3, 5, 10 or more residues, from a sequence of the invention contained in the Sequence Listing. The differences, however, are such that: the *M. catarrhalis* encoded polypeptide exhibits an *M. catarrhalis* biological activity, e.g., the encoded *M. catarrhalis* enzyme retains a biological activity of a naturally occurring *M. catarrhalis*.

In preferred embodiments, the encoded polypeptide includes all or a fragment of an amino acid sequence of the invention contained in the Sequence Listing; fused, in reading

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frame, to additional amino acid residues, preferably to residues encoded by genomic DNA 5' or 3' to the genomic DNA which encodes a sequence of the invention contained in the Sequence Listing.

The *M. catarrhalis* strain, 98-4362, from which genomic sequences have been sequenced, has been deposited on July 20, 1998, in the American Type Culture Collection and assigned the ATCC designation # 202156.

Included in the invention are: allelic variations; natural mutants; induced mutants; proteins encoded by DNA that hybridize under high or low stringency conditions to a nucleic acid which encodes a polypeptide of the invention contained in the Sequence Listing (for definitions of high and low stringency see Current Protocols in Molecular Biology, John Wiley & Sons, New York, 1989, 6.3.1 - 6.3.6, hereby incorporated by reference); and, polypeptides specifically bound by antisera to *M. catarrhalis* polypeptides, especially by antisera to an active site or binding domain of *M. catarrhalis* polypeptide. The invention also includes fragments, preferably biologically active fragments. These and other polypeptides are also referred to herein as *M. catarrhalis* polypeptide analogs or variants.

The invention further provides nucleic acids, e.g., RNA or DNA and their respective complements, encoding a polypeptide of the invention. This includes double stranded nucleic acids as well as coding and antisense single strands.

In preferred embodiments, the subject *M. catarrhalis* nucleic acid will include a transcriptional regulatory sequence, e.g., at least one of a transcriptional promoter or transcriptional enhancer sequence, operably linked to the *M. catarrhalis* gene sequence, e.g., to render the *M. catarrhalis* gene sequence suitable for expression in a recombinant host cell.

In yet a further preferred embodiment, the nucleic acid which encodes an M. catarrhalis polypeptide of the invention, hybridizes under stringent conditions to a nucleic acid probe corresponding to at least about 8 consecutive nucleotides of the invention contained in the Sequence Listing; more preferably to at least about 12 consecutive nucleotides of the invention contained in the Sequence Listing; still more preferably to at least about 20 consecutive nucleotides of the invention contained in the Sequence Listing; most preferably to at least about 40 consecutive nucleotides of the invention contained in the Sequence Listing.

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In another aspect, the invention provides a substantially pure nucleic acid having a nucleotide sequence which encodes an *M. catarrhalis* polypeptide. In preferred embodiments: the encoded polypeptide has biological activity; the encoded polypeptide has an amino acid sequence at least about 60%, 70%, 80%, 90%, 95%, 98% or 99% homologous to an amino acid sequence of the invention contained in the Sequence Listing; the encoded polypeptide has an amino acid sequence essentially the same as an amino acid sequence of the invention contained in the Sequence Listing; the encoded polypeptide is at least about 5, 10, 20, 50, 100, or 150 amino acids in length; the encoded polypeptide comprises at least about 5, preferably at least about 10, more preferably at least about 20, still more preferably at least about 50, 100, or 150 contiguous amino acids of the invention contained in the Sequence Listing.

In another aspect, the invention encompasses: a vector including a nucleic acid which encodes an *M. catarrhalis* polypeptide or an *M. catarrhalis* polypeptide variant as described herein; a host cell transfected with the vector; and a method of producing a recombinant *M. catarrhalis* polypeptide or *M. catarrhalis* polypeptide variant; including culturing the cell, e.g., in a cell culture medium, and isolating *an M. catarrhalis* or *M. catarrhalis* polypeptide variant, e.g., from the cell or from the cell culture medium.

One embodiment of the invention is directed to substantially isolated nucleic acids. Nucleic acids of the invention include sequences comprising at least about 8 nucleotides in length, more preferably at least about 12 nucleotides in length, even more preferably at least about 15-20 nucleotides in length, that correspond to a subsequence of any one of SEQ ID NO: 1 - SEQ ID NO: 1920 or complements thereof. Alternatively, the nucleic acids comprise sequences contained within any ORF (open reading frame), including a complete protein-coding sequence, of which any of SEQ ID NO: 1 - SEQ ID NO: 1920 forms a part. The invention encompasses sequence-conservative variants and function-conservative variants of these sequences. The nucleic acids may be DNA, RNA, DNA/RNA duplexes, protein-nucleic acid (PNA), or derivatives thereof.

In another aspect, the invention features a purified recombinant nucleic acid having at least about 50%, 60%, 70%, 80%, 90%, 95%, 98%, or 99% homology with a sequence of the invention contained in the Sequence Listing

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The invention also encompasses recombinant DNA (including DNA cloning and expression vectors) comprising these *M. catarrhalis* -derived sequences; host cells comprising such DNA, including fungal, bacterial, yeast, plant, insect, and mammalian host cells; and methods for producing expression products comprising RNA and polypeptides encoded by the *M. catarrhalis* sequences. These methods are carried out by incubating a host cell comprising an *M. catarrhalis* -derived nucleic acid sequence under conditions in which the sequence is expressed. The host cell may be native or recombinant. The polypeptides can be obtained by (a) harvesting the incubated cells to produce a cell fraction and a medium fraction; and (b) recovering the *M. catarrhalis* polypeptide from the cell fraction, the medium fraction, or both. The polypeptides can also be made by *in vitro* translation.

In another aspect, the invention features nucleic acids capable of binding mRNA of *M. catarrhalis*. Such nucleic acid is capable of acting as antisense nucleic acid to control the translation of mRNA of *M. catarrhalis*. A further aspect features a nucleic acid which is capable of binding specifically to an *M. catarrhalis* nucleic acid. These nucleic acids are also referred to herein as complements and have utility as probes and as capture reagents.

In another aspect, the invention features an expression system comprising an open reading frame corresponding to *M. catarrhalis* nucleic acid. The nucleic acid further comprises a control sequence compatible with an intended host. The expression system is useful for making polypeptides corresponding to *M. catarrhalis* nucleic acid.

In another aspect, the invention encompasses: a vector including a nucleic acid which encodes an *M. catarrhalis* polypeptide or an *M. catarrhalis* polypeptide variant as described herein; a host cell transfected with the vector; and a method of producing a recombinant *M. catarrhalis* polypeptide or *M. catarrhalis* polypeptide variant; including culturing the cell, e.g., in a cell culture medium, and isolating the *M. catarrhalis* or *M. catarrhalis* polypeptide variant, e.g., from the cell or from the cell culture medium.

In yet another embodiment of the invention encompasses reagents for detecting bacterial infection, including *M. catarrhalis* infection, which comprise at least one *M. catarrhalis* -derived nucleic acid defined by any one of SEQ ID NO: 1 - SEQ ID NO: 1920, or sequence-conservative or function-conservative variants thereof. Alternatively, the

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diagnostic reagents comprise nucleotide sequences that are contained within any open reading frames (ORFs), including preferably complete protein-coding sequences, contained within any of SEQ ID NO: 1 - SEQ ID NO: 1920, or polypeptide sequences contained within any of SEQ ID NO: 1921 - SEQ ID NO: 3840, or polypeptides of which any of the above sequences forms a part, or antibodies directed against any of the above peptide sequences or function-conservative variants and/or fragments thereof.

The invention further provides antibodies, preferably monoclonal antibodies, which specifically bind to the polypeptides of the invention. Methods are also provided for producing antibodies in a host animal. The methods of the invention comprise immunizing an animal with at least one *M. catarrhalis* -derived immunogenic component, wherein the immunogenic component comprises one or more of the polypeptides encoded by any one of SEQ ID NO: 1 - SEQ ID NO: 1920 or sequence-conservative or function-conservative variants thereof; or polypeptides that are contained within any ORFs, including complete protein-coding sequences, of which any of SEQ ID NO: 1 - SEQ ID NO: 1920 forms a part; or polypeptide sequences contained within any of SEQ ID NO: 1921 - SEQ ID NO: 3840; or polypeptides of which any of SEQ ID NO: 1921 - SEQ ID NO: 3840 forms a part. Host animals include any warm blooded animal, including without limitation mammals and birds. Such antibodies have utility as reagents for immunoassays to evaluate the abundance and distribution of *M. catarrhalis* -specific antigens.

In yet another aspect, the invention provides diagnostic methods for detecting *M. catarrhalis* antigenic components or anti-*M. catarrhalis* antibodies in a sample. *M. catarrhalis* antigenic components may be detected by known processes, including but not limited to detection by a process comprising: (i) contacting a sample suspected to contain a bacterial antigenic component with a bacterial-specific antibody, under conditions in which a stable antigen-antibody complex can form between the antibody and bacterial antigenic components in the sample; and (ii) detecting any antigen-antibody complex formed in step (i), wherein detection of an antigen-antibody complex indicates the presence of at least one bacterial antigenic component in the sample. In different embodiments of this method, the antibodies used are directed against a sequence encoded by any of SEQ ID NO: 1 - SEQ ID NO: 1920 or sequence-conservative or function-conservative variants thereof, or against a

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polypeptide sequence contained in any of SEQ ID NO: 1921 - SEQ ID NO: 3840 or function-conservative variants thereof.

In yet another aspect, the invention provides a method for detecting antibacterial-specific antibodies in a sample, which comprises: (i) contacting a sample suspected to contain antibacterial-specific antibodies with an *M. catarrhalis* antigenic component, under conditions in which a stable antigen-antibody complex can form between the *M. catarrhalis* antigenic component and antibacterial antibodies in the sample; and (ii) detecting any antigen-antibody complex formed in step (i), wherein detection of an antigen-antibody complex indicates the presence of antibacterial antibodies in the sample. In different embodiments of this method, the antigenic component is encoded by a sequence contained in any of SEQ ID NO: 1 - SEQ ID NO: 1920 or sequence-conservative and function-conservative variants thereof, or is a polypeptide sequence contained in any of SEQ ID NO: 1921 - SEQ ID NO: 3840 or function-conservative variants thereof.

In another aspect, the invention features a method of generating vaccines for immunizing an individual against *M. catarrhalis*. The method includes: immunizing a subject with an *M. catarrhalis* polypeptide, e.g., a surface or secreted polypeptide, or a combination of such peptides or active portion(s) thereof, and a pharmaceutically acceptable carrier. Such vaccines have therapeutic and prophylactic utilities.

In another aspect, the invention features a method of evaluating a compound, e.g., a polypeptide, e.g., a fragment of a host cell polypeptide, for the ability to bind an M. catarrhalis polypeptide. The method includes contacting the compound to be evaluated with an M. catarrhalis polypeptide and determining if the compound binds or otherwise interacts with the M catarrhalis polypeptide. Compounds which bind or otherwise interact with M catarrhalis polypeptides are candidates as modulators, including activators and inhibitors, of the bacterial life cycle. These assays can be performed in vitro or in vivo.

In another aspect, the invention features a method of evaluating a compound, e.g., a polypeptide, e.g., a fragment of a host cell polypeptide, for the ability to bind an M. catarrhalis nucleic acid, e.g., DNA or RNA. The method includes contacting the compound to be evaluated with an M. catarrhalis nucleic acid and determining if the compound binds or otherwise interacts with the M catarrhalis nucleic acid. Compounds which bind M.

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catarrhalis are candidates as modulators, including activators and inhibitors, of the bacterial life cycle. These assays can be performed in vitro or in vivo.

A particularly preferred embodiment of the invention is directed to a method of screening test compounds for anti-bacterial activity, which method comprises: selecting as a target a bacterial specific sequence, which sequence is essential to the viability of a bacterial species; contacting a test compound with said target sequence; and selecting those test compounds which bind to said target sequence as potential anti-bacterial candidates. In one embodiment, the target sequence selected is specific to a single species, or even a single strain, such as, for example, the strain *M. catarrhalis*98-4362. In a second embodiment, the target sequence is common to at least two species of bacteria. In a third embodiment, the target sequence is common to a family of bacteria. The target sequence may be a nucleic acid sequence or a polypeptide sequence. Methods employing sequences common to more than one species of microorganism may be used to screen candidates for broad spectrum anti-bacterial activity.

The invention also provides methods for preventing or treating disease caused by certain bacteria, including *M. catarrhalis*, which are carried out by administering to an animal in need of such treatment, in particular a warm-blooded vertebrate, including but not limited to birds and mammals, a compound that specifically inhibits or interferes with the function of a bacterial polypeptide or nucleic acid. In a particularly preferred embodiment, the mammal to be treated is human.

#### DETAILED DESCRIPTION OF THE INVENTION

The sequences of the present invention include the specific nucleic acid and amino acid sequences set forth in the Sequence Listing that forms a part of the present specification, and which are designated SEQ ID NO: 1 - SEQ ID NO: 3840. Use of the terms "SEQ ID NO: 1 - SEQ ID NO: 1920 ", " SEQ ID NO: 1921 - SEQ ID NO: 3840, "the sequences depicted in Table 2", etc., is intended, for convenience, to refer to each individual SEQ ID NO individually, and is not intended to refer to the genus of these sequences unless such reference would be indicated. In other words, it is a shorthand for listing all of these

sequences individually. The invention encompasses each sequence individually, as well as any combination thereof.

#### **DEFINITIONS**

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"Nucleic acid" or "polynucleotide" as used herein refers to purine- and pyrimidine-containing polymers of any length, either polyribonucleotides or polydeoxyribonucleotides or mixed polyribo-polydeoxyribo nucleotides. This includes single- and double-stranded molecules, i.e., DNA-DNA, DNA-RNA and RNA-RNA hybrids, as well as "protein nucleic acids" (PNA) formed by conjugating bases to an amino acid backbone. This also includes nucleic acids containing modified bases.

A nucleic acid or polypeptide sequence that is "derived from" a designated sequence refers to a sequence that corresponds to a region of the designated sequence. For nucleic acid sequences, this encompasses sequences that are homologous or complementary to the sequence, as well as "sequence-conservative variants" and "function-conservative variants." For polypeptide sequences, this encompasses "function-conservative variants." Sequence-conservative variants are those in which a change of one or more nucleotides in a given codon position results in no alteration in the amino acid encoded at that position. Function-conservative variants are those in which a given amino acid residue in a polypeptide has been changed without altering the overall conformation and function of the native polypeptide, including, but not limited to, replacement of an amino acid with one having similar physico-chemical properties (such as, for example, acidic, basic, hydrophobic, and the like). "Function-conservative" variants also include any polypeptides that have the ability to elicit antibodies specific to a designated polypeptide.

An "M. catarrhalis -derived" nucleic acid or polypeptide sequence may or may not be present in other bacterial species, and may or may not be present in all M. catarrhalis strains. This term is intended to refer to the source from which the sequence was originally isolated. Thus, an M. catarrhalis -derived polypeptide, as used herein, may be used, e.g., as a target to screen for a broad spectrum antibacterial agent, to search for homologous proteins in other species of bacteria or in eukaryotic organisms such as bacteria humans, etc.

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A purified or isolated polypeptide or a substantially pure preparation of a polypeptide are used interchangeably herein and, as used herein, mean a polypeptide that has been separated from other proteins, lipids, and nucleic acids with which it naturally occurs. Preferably, the polypeptide is also separated from substances, e.g., antibodies or gel matrix, e.g., polyacrylamide, which are used to purify it. Preferably, the polypeptide constitutes at least about 10, 20, 50 70, 80 or 95% dry weight of the purified preparation. Preferably, the preparation contains sufficient polypeptide to allow protein sequencing; at least about 1, 10, or preferably 100 mg of polypeptide.

A purified preparation of cells refers to, in the case of plant or animal cells, an *in* vitro preparation of cells and not an entire intact plant or animal. In the case of cultured cells or microbial cells, it consists of a preparation of at least about 10%, more preferably at least about 50%, of the subject cells.

A purified or isolated or a substantially pure nucleic acid, e.g., a substantially pure DNA, (are terms used interchangeably herein) is a nucleic acid which is one or both of the following: not immediately contiguous with both of the coding sequences with which it is immediately contiguous (i.e., one at the 5' end and one at the 3' end) in the naturally-occurring genome and plasmids of the organism from which the nucleic acid is derived; or which is substantially free of a nucleic acid with which it occurs in the organism from which the nucleic acid is derived. The term includes, for example, a recombinant DNA which is incorporated into a vector, e.g., into an autonomously replicating plasmid or virus, or into the genomic DNA of a prokaryote or eukaryote, or which exists as a separate molecule (e.g., a cDNA or a genomic DNA fragment produced by PCR or restriction endonuclease treatment) independent of other DNA sequences. Substantially pure DNA also includes a recombinant DNA which is part of a hybrid gene encoding additional *M. catarrhalis* DNA sequence.

A "contig" as used herein is a nucleic acid representing a continuous stretch of genomic sequence of an organism.

An "open reading frame", also referred to herein as ORF, is a region of nucleic acid which encodes a polypeptide. This region may represent a portion of a coding sequence or a total sequence and can be determined from a stop to stop codon or from a start to stop codon.

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As used herein, a "coding sequence" is a nucleic acid which is transcribed into messenger RNA and/or translated into a polypeptide when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a translation start codon at the five prime terminus and a translation stop code at the three prime terminus. A coding sequence can include but is not limited to messenger RNA, synthetic DNA, and recombinant nucleic acid sequences.

A "complement" of a nucleic acid as used herein refers to an anti-parallel or antisense sequence that participates in Watson-Crick base-pairing with the original sequence.

A "gene product" is a protein or structural RNA which is specifically encoded by a gene.

As used herein, the term "probe" refers to a nucleic acid, peptide or other chemical entity which specifically binds to a molecule of interest. Probes are often associated with or capable of associating with a label. A label is a chemical moiety capable of detection.

Typical labels comprise dyes, radioisotopes, luminescent and chemiluminescent moieties, fluorophores, enzymes, precipitating agents, amplification sequences, and the like.

Similarly, a nucleic acid, peptide or other chemical entity which specifically binds to a molecule of interest and immobilizes such molecule is referred herein as a "capture ligand".

Capture ligands are typically associated with or capable of associating with a support such as nitro-cellulose, glass, nylon membranes, beads, particles and the like. The specificity of hybridization is dependent on conditions such as the base pair composition of the nucleotides, and the temperature and salt concentration of the reaction. These conditions are readily discernable to one of ordinary skill in the art using routine experimentation.

"Homologous" refers to the sequence similarity or sequence identity between two polypeptides or between two nucleic acid molecules. When a position in both of the two compared sequences is occupied by the same base or amino acid monomer subunit, e.g., if a position in each of two DNA molecules is occupied by adenine, then the molecules are homologous at that position. The percent of homology between two sequences is a function of the number of matching or homologous positions shared by the two sequences divided by the number of positions compared x 100. For example, if 6 of 10 of the positions in two sequences are matched or homologous then the two sequences are 60% homologous. By

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way of example, the DNA sequences ATTGCC and TATGGC share 50% homology. Generally, a comparison is made when two sequences are aligned to give maximum homology.

Nucleic acids are hybridizable to each other when at least one strand of a nucleic acid can anneal to the other nucleic acid under defined stringency conditions. Stringency of 5 hybridization is determined by: (a) the temperature at which hybridization and/or washing is performed; and (b) the ionic strength and polarity of the hybridization and washing solutions. Hybridization requires that the two nucleic acids contain complementary sequences; depending on the stringency of hybridization, however, mismatches may be tolerated. Typically, hybridization of two sequences at high stringency (such as, for example, in a 10 solution of 0.5X SSC, at 65° C) requires that the sequences be essentially completely homologous. Conditions of intermediate stringency (such as, for example, 2X SSC at 65° C) and low stringency (such as, for example 2X SSC at 55°C) require correspondingly less overall complementarity between the hybridizing sequences. (1X SSC is 0.15 M NaCl, 0.015 M Na citrate).

The terms peptides, proteins, and polypeptides are used interchangeably herein.

As used herein, the term "surface protein" refers to all surface accessible proteins, e.g. inner and outer membrane proteins, proteins adhering to the cell wall, and secreted proteins.

A polypeptide has M. catarrhalis biological activity if it has one, two or preferably more of the following properties: (1) if when expressed in the course of an M. catarrhalis infection, it can promote, or mediate the attachment of M. catarrhalis to a cell; (2) it has an enzymatic activity, structural or regulatory function characteristic of an M. catarrhalis protein; (3) the gene which encodes it can rescue a lethal mutation in an M. catarrhalis gene. A polypeptide has biological activity if it is an antagonist, agonist, or super-agonist of a polypeptide having one of the above-listed properties.

A biologically active fragment or analog is one having an in vivo or in vitro activity which is characteristic of the M. catarrhalis polypeptides of the invention contained in the Sequence Listing, or of other naturally occurring M. catarrhalis polypeptides, e.g., one or more of the biological activities described herein. Especially preferred are fragments which

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exist *in vivo*, e.g., fragments which arise from post transcriptional processing or which arise from translation of alternatively spliced RNA's. Fragments include those expressed in native or endogenous cells as well as those made in expression systems, e.g., in CHO (Chinese Hamster Ovary) cells. Because peptides such as *M. catarrhalis* polypeptides often exhibit a range of physiological properties and because such properties may be attributable to different portions of the molecule, a useful *M. catarrhalis* fragment or *M. catarrhalis* analog is one which exhibits a biological activity in any biological assay for *M. catarrhalis* activity. The fragment or analog possesses about 10%, preferably about 40%, more preferably about 60%, 70%, 80% or 90% or greater of the activity of *M. catarrhalis*, in any *in vivo* or *in vitro* assay.

Analogs can differ from naturally occurring *M. catarrhalis* polypeptides in amino acid sequence or in ways that do not involve sequence, or both. Non-sequence modifications include changes in acetylation, methylation, phosphorylation, carboxylation, or glycosylation. Preferred analogs include *M. catarrhalis* polypeptides (or biologically active fragments thereof) whose sequences differ from the wild-type sequence by one or more conservative amino acid substitutions or by one or more non-conservative amino acid substitutions, deletions, or insertions which do not substantially diminish the biological activity of the *M. catarrhalis* polypeptide. Conservative substitutions typically include the substitution of one amino acid for another with similar characteristics, e.g., substitutions within the following groups: valine, glycine; glycine, alanine; valine, isoleucine, leucine; aspartic acid, glutamic acid; asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, tyrosine. Other conservative substitutions can be made in view of the table below.

25 TABLE 1
CONSERVATIVE AMINO ACID REPLACEMENTS

For Amino Acid	Code	Replace with any of
Alanine	A	D-Ala, Gly, beta-Ala, L-Cys, D-Cys

Arginine	R	D-Arg, Lys, D-Lys, homo-Arg, D-homo-Arg, Met, Ile, D-Met, D-Ile, Orn, D-Orn
Asparagine	N	D-Asn, Asp, D-Asp, Glu, D-Glu, Gln, D-Gln
Aspartic Acid	D .	D-Asp, D-Asn, Asn, Glu, D-Glu, Gln, D-Gln
Cysteine	С	D-Cys, S-Me-Cys, Met, D-Met, Thr, D-Thr
Glutamine	Q	D-Gln, Asn, D-Asn, Glu, D-Glu, Asp, D-Asp
Glutamic Acid	E	D-Glu, D-Asp, Asp, Asn, D-Asn, Gln, D-Gln
Glycine	G	Ala, D-Ala, Pro, D-Pro, β-Ala, Acp
Isoleucine	I	D-Ile, Val, D-Val, Leu, D-Leu, Met, D-Met
Leucine	L .	D-Leu, Val, D-Val, Leu, D-Leu, Met, D-Met
Lysine	K	D-Lys, Arg, D-Arg, homo-Arg, D-homo-Arg, Met, D-Met, Ile, D-Ile, Orn, D-Orn
Methionine	М	D-Met, S-Me-Cys, Ile, D-Ile, Leu, D-Leu, Val, D-Val
Phenylalanine	F	D-Phe, Tyr, D-Thr, L-Dopa, His, D-His, Trp, D-Trp, Trans-3,4, or 5-phenylproline, cis-3,4, or 5-phenylproline
Proline	P	D-Pro, L-I-thioazolidine-4-carboxylic acid, D-or L-1-oxazolidine-4-carboxylic acid
Serine	S	D-Ser, Thr, D-Thr, allo-Thr, Met, D-Met, Met(O), D-Met(O), L-Cys, D-Cys
Threonine	T	D-Thr, Ser, D-Ser, allo-Thr, Met, D-Met, Met(O), D-Met(O), Val, D-Val
Tyrosine	Y	D-Tyr, Phe, D-Phe, L-Dopa, His, D-His
Valine	V	D-Val, Leu, D-Leu, Ile, D-Ile, Met, D-Met

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Other analogs within the invention are those with modifications which increase peptide stability; such analogs may contain, for example, one or more non-peptide bonds (which replace the peptide bonds) in the peptide sequence. Also included are: analogs that include residues other than naturally occurring L-amino acids, e.g., D-amino acids or non-naturally occurring or synthetic amino acids, e.g.,  $\beta$  or  $\gamma$  amino acids; and cyclic analogs.

As used herein, the term "fragment", as applied to an *M. catarrhalis* analog, will ordinarily be at least about 20 residues, more typically at least about 40 residues, preferably at least about 60 residues in length. Fragments of *M. catarrhalis* polypeptides can be generated by methods known to those skilled in the art. The ability of an *Moraxella* fragment to exhibit a biological activity of *M. catarrhalis* polypeptide can be assessed by methods known to those skilled in the art as described herein. Also included are *M. catarrhalis* polypeptides containing residues that are not required for biological activity of the peptide or that result from alternative mRNA splicing or alternative protein processing events.

An "immunogenic component" as used herein is a moiety, such as an *M. catarrhalis* polypeptide, analog or fragment thereof, that is capable of eliciting a humoral and/or cellular immune response in a host animal.

An "antigenic component" as used herein is a moiety, such as an *M. catarrhalis* polypeptide, analog or fragment thereof, that is capable of binding to a specific antibody with sufficiently high affinity to form a detectable antigen-antibody complex.

The term "antibody" as used herein is intended to include fragments thereof which are specifically reactive with *M. catarrhalis* polypeptides.

As used herein, the term "cell-specific promoter" means a DNA sequence that serves as a promoter, i.e., regulates expression of a selected DNA sequence operably linked to the promoter, and which effects expression of the selected DNA sequence in specific cells of a tissue. The term also covers so-called "leaky" promoters, which regulate expression of a selected DNA primarily in one tissue, but cause expression in other tissues as well.

Misexpression, as used herein, refers to a non-wild type pattern of gene expression. It includes: expression at non-wild type levels, i.e., over or under expression; a pattern of expression that differs from wild type in terms of the time or stage at which the gene is

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expressed, e.g., increased or decreased expression (as compared with wild type) at a predetermined developmental period or stage; a pattern of expression that differs from wild type in terms of increased expression (as compared with wild type) in a predetermined cell type or tissue type; a pattern of expression that differs from wild type in terms of the splicing size, amino acid sequence, post-translational modification, or biological activity of the expressed polypeptide; a pattern of expression that differs from wild type in terms of the effect of an environmental stimulus or extracellular stimulus on expression of the gene, e.g., a pattern of increased or decreased expression (as compared with wild type) in the presence of an increase or decrease in the strength of the stimulus.

As used herein, "host cells" and other such terms denoting microorganisms or higher eukaryotic cell lines cultured as unicellular entities refers to cells which can become or have been used as recipients for a recombinant vector or other transfer DNA, and include the progeny of the original cell which has been transfected. It is understood by individuals skilled in the art that the progeny of a single parental cell may not necessarily be completely identical in genomic or total DNA compliment to the original parent, due to accident or deliberate mutation.

As used herein, the term "control sequence" refers to a nucleic acid having a base sequence which is recognized by the host organism to effect the expression of encoded sequences to which they are ligated. The nature of such control sequences differs depending upon the host organism; in prokaryotes, such control sequences generally include a promoter, ribosomal binding site, terminators, and in some cases operators; in eukaryotes, generally such control sequences include promoters, terminators and in some instances, enhancers. The term control sequence is intended to include at a minimum, all components whose presence is necessary for expression, and may also include additional components whose presence is advantageous, for example, leader sequences.

As used herein, the term "operably linked" refers to sequences joined or ligated to function in their intended manner. For example, a control sequence is operably linked to coding sequence by ligation in such a way that expression of the coding sequence is achieved under conditions compatible with the control sequence and host cell.

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The "metabolism" of a substance, as used herein, means any aspect of the expression, function, action, or regulation of the substance. The metabolism of a substance includes modifications, e.g., covalent or non-covalent modifications of the substance. The metabolism of a substance includes modifications, e.g., covalent or non-covalent modification, the substance induces in other substances. The metabolism of a substance also includes changes in the distribution of the substance. The metabolism of a substance includes changes the substance induces in the distribution of other substances.

A "sample" as used herein refers to a biological sample, such as, for example, tissue or fluid isloated from an individual (including without limitation plasma, serum, cerebrospinal fluid, lymph, tears, saliva and tissue sections) or from *in vitro* cell culture constituents, as well as samples from the environment.

Technical and scientific terms used herein have the meanings commonly understood by one of ordinary skill in the art to which the present invention pertains, unless otherwise defined. Reference is made herein to various methodologies known to those of skill in the art. Publications and other materials setting forth such known methodologies to which reference is made are incorporated herein by reference in their entireties as though set forth in full. The practice of the invention will employ, unless otherwise indicated, conventional techniques of chemistry, molecular biology, microbiology, recombinant DNA, and immunology, which are within the skill of the art. Such techniques are explained fully in the literature. See e.g., Sambrook, Fritsch, and Maniatis, Molecular Cloning: Laboratory Manual 2nd ed. (1989); DNA Cloning, Volumes I and II (D.N Glover ed. 1985); Oligonucleotide Synthesis (M.J. Gait ed, 1984); Nucleic Acid Hybridization (B.D. Hames & S.J. Higgins eds. 1984); the series, Methods in Enzymology (Academic Press, Inc.), particularly Vol. 154 and Vol. 155 (Wu and Grossman, eds.); PCR-A Practical Approach (McPherson, Quirke, and Taylor, eds., 1991); Immunology, 2d Edition, 1989, Roitt et al., C.V. Mosby Company, and New York; Advanced Immunology, 2d Edition, 1991, Male et al., Grower Medical Publishing, New York.; DNA Cloning: A Practical Approach, Volumes I and II, 1985 (D.N. Glover ed.); Oligonucleotide Synthesis, 1984, (M.L. Gait ed); Transcription and Translation, 1984 (Hames and Higgins eds.); Animal Cell Culture, 1986 (R.I. Freshney ed.); Immobilized Cells and Enzymes, 1986 (IRL Press); Perbal, 1984, A

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Practical Guide to Molecular Cloning; Gene Transfer Vectors for Mammalian Cells, 1987 (J. H. Miller and M. P. Calos eds., Cold Spring Harbor Laboratory); Martin J. Bishop, ed., Guide to Human Genome Computing, 2d Edition, Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, The Internet and the New Biology: Tools for Genomic and Molecular Research, American Society for Microbiology, Washington, D.C. (1997).

Any suitable materials and/or methods known to those of skill can be utilized in carrying out the present invention; however, preferred materials and/or methods are described. Materials, reagents and the like to which reference is made in the following description and examples are obtainable from commercial sources, unless otherwise noted.

#### M. CATARRHALIS GENOMIC SEQUENCE

This invention provides nucleotide sequences of the genome of *M. catarrhalis* which thus comprises a DNA sequence library of *M. catarrhalis* genomic DNA. The detailed description that follows provides nucleotide sequences of *M. catarrhalis*, and also describes how the sequences were obtained and how ORFs and protein-coding sequences were identified. Also described are compositions and methods of using the disclosed *M. catarrhalis* sequences in methods including diagnostic and therapeutic applications. Furthermore, the library can be used as a database for identification and comparison of medically important sequences in this and other strains of *M. catarrhalis*.

To determine the genomic sequence of *M. catarrhalis*, DNA from strain 98-4362. of *M. catarrhalis* was isolated and a library of DNA fragments were transformed into DH5α cells. DNA sequencing was achieved using established ABI sequencing methods on ABI377 automated DNA sequencers. The cloning and sequencing procedures are described in more detail in the Exemplification.

Individual sequence reads were assembled using PHRAP (P. Green, Abstracts of DOE Human Genome Program Contractor-Grantee Workshop V, Jan. 1996, p.157). The average contig length was about 3-4 kb.

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All subsequent steps were based on sequencing by ABI377 automated DNA sequencing methods. The cloning and sequencing procedures are described in more detail in the Exemplification.

A variety of approaches may be used to order the contigs so as to obtain a continuous sequence representing the entire *M. catarrhalis* genome. Synthetic oligonucleotides are designed that are complementary to sequences at the end of each contig. These oligonucleotides may be hybridized to libaries of *M. catarrhalis* genomic DNA in, for example, lambda phage vectors or plasmid vectors to identify clones that contain sequences corresponding to the junctional regions between individual contigs. Such clones are then used to isolate template DNA and the same oligonucleotides are used as primers in polymerase chain reaction (PCR) to amplify junctional fragments, the nucleotide sequence of which is then determined.

The M catarrhalis sequences were analyzed for the presence of open reading frames (ORFs) comprising at least 180 nucleotides. As a result of the analysis of ORFs based on stop-to-stop codon reads, it should be understood that these ORFs may not correspond to the ORF of a naturally-occurring M catarrhalis polypeptide. These ORFs may contain start codons which indicate the initiation of protein synthesis of a naturally-occurring M catarrhalis polypeptide. Such start codons within the ORFs provided herein were identified by those of ordinary skill in the relevant art, and the resulting ORF and the encoded M catarrhalis polypeptide is within the scope of this invention. For example, within the ORFs a codon such as AUG or GUG (encoding methionine or valine) which is part of the initiation signal for protein synthesis were identified and the portion of an ORF to corresponding to a naturally-occurring M catarrhalis polypeptide was recognized. The predicted coding regions were defined by evaluating the coding potential of such sequences with the program GENEMARK<sup>TM</sup> (Borodovsky and McIninch, 1993, Comp. 17:123).

Each predicted ORF amino acid sequence was compared with all sequences found in current GENBANK, SWISS-PROT, and PIR databases using the BLAST algorithm. BLAST identifies local alignments occurring by chance between the ORF sequence and the sequence in the databank (Altschal et al., 1990, L Mol. Biol. 215:403-410). Homologous ORFs (probabilities less than 10<sup>-5</sup> by chance) and ORF's that are probably non-homologous

(probabilities greater than 10<sup>-5</sup> by chance) but have good codon usage were identified. Both homologous, sequences and non-homologous sequences with good codon usage, are likely to encode proteins and are encompassed by the invention.

#### 5 M. CATARRHALIS NUCLEIC ACIDS

The present invention provides a library of *M. catarrhalis* -derived nucleic acid sequences. The libraries provide probes, primers, and markers which are used as markers in epidemiological studies. The present invention also provides a library of *M. catarrhalis* - derived nucleic acid sequences which comprise or encode targets for therapeutic drugs.

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The nucleic acids of this invention may be obtained directly from the DNA of the above referenced *M. catarrhalis* strain by using the polymerase chain reaction (PCR). See "PCR, A Practical Approach" (McPherson, Quirke, and Taylor, eds., IRL Press, Oxford, UK, 1991) for details about the PCR. High fidelity PCRis used to ensure a faithful DNA copy prior to expression. In addition, the authenticity of amplified products is verified by conventional sequencing methods. Clones carrying the desired sequences described in this invention may also be obtained by screening the libraries by means of the PCR or by hybridization of synthetic oligonucleotide probes to filter lifts of the library colonies or plaques as known in the art (see, e.g., Sambrook et al., Molecular Cloning, A Laboratory Manual 2nd edition, 1989, Cold Spring Harbor Press, NY).

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It is also possible to obtain nucleic acids encoding *M. catarrhalis* polypeptides from a cDNA library in accordance with protocols herein described. A cDNA encoding an *M. catarrhalis* polypeptide can be obtained by isolating total mRNA from an appropriate strain. Double stranded cDNAs can then be prepared from the total mRNA. Subsequently, the cDNAs can be inserted into a suitable plasmid or viral (e.g., bacteriophage) vector using any one of a number of known techniques. Genes encoding *M. catarrhalis* polypeptides can also be cloned using established polymerase chain reaction techniques in accordance with the nucleotide sequence information provided by the invention. The nucleic acids of the invention can be DNA or RNA. Preferred nucleic acids of the invention are contained in the Sequence Listing.

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The nucleic acids of the invention can also be chemically synthesized using standard techniques. Various methods of chemically synthesizing polydeoxynucleotides are known, including solid-phase synthesis which, like peptide synthesis, has been fully automated in commercially available DNA synthesizers (See e.g., Itakura et al. U.S. Patent No. 4,598,049; Caruthers et al. U.S. Patent No. 4,458,066; and Itakura U.S. Patent Nos. 4,401,796 and 4,373,071, incorporated by reference herein).

In another example, DNA can be chemically synthesized using, e.g., the phosphoramidite solid support method of Matteucci *et al.*, 1981, *J. Am. Chem. Soc.* 103:3185, the method of Yoo *et al.*, 1989, *J. Biol. Chem.* 764:17078, or other well known methods. This can be done by sequentially linking a series of oligonucleotide cassettes comprising pairs of synthetic oligonucleotides, as described below.

Nucleic acids isolated or synthesized in accordance with features of the present invention are useful, by way of example, without limitation, as probes, primers, capture ligands, antisense genes and for developing expression systems for the synthesis of proteins and peptides corresponding to such sequences. As probes, primers, capture ligands and antisense agents, the nucleic acid normally consists of all or part (approximately twenty or more nucleotides for specificity as well as the ability to form stable hybridization products) of the nucleic acids of the invention contained in the Sequence Listing. These uses are described in further detail below.

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#### **PROBES**

A nucleic acid isolated or synthesized in accordance with the sequence of the invention contained in the Sequence Listing can be used as a probe to specifically detect *M. catarrhalis*. With the sequence information set forth in the present application, sequences of twenty or more nucleotides are identified which provide the desired inclusivity and exclusivity with respect to *M. catarrhalis*, and extraneous nucleic acids likely to be encountered during hybridization conditions. More preferably, the sequence will comprise at least about twenty to thirty nucleotides to convey stability to the hybridization product formed between the probe and the intended target molecules.

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Sequences larger than 1000 nucleotides in length are difficult to synthesize but can be generated by recombinant DNA techniques. Individuals skilled in the art will readily recognize that the nucleic acids, for use as probes, can be provided with a label to facilitate detection of a hybridization product.

Nucleic acid isolated and synthesized in accordance with the sequence of the invention contained in the Sequence Listing can also be useful as probes to detect homologous regions (especially homologous genes) of other *Moraxella* species using appropriate stringency hybridization conditions as described herein.

#### 10 CAPTURE LIGAND

For use as a capture ligand, the nucleic acid selected in the manner described above with respect to probes, can be readily associated with a support. The manner in which nucleic acid is associated with supports is well known. Nucleic acid having twenty or more nucleotides in a sequence of the invention contained in the Sequence Listing have utility to separate *M. catarrhalis* nucleic acid from one strain from the nucleic acid of other another strain as well as from other organisms. Nucleic acid having twenty or more nucleotides in a sequence of the invention contained in the Sequence Listing can also have utility to separate other *Moraxella* species from each other and from other organisms. Preferably, the sequence will comprise at least about twenty nucleotides to convey stability to the hybridization product formed between the probe and the intended target molecules. Sequences larger than 1000 nucleotides in length are difficult to synthesize but can be generated by recombinant DNA techniques.

#### **PRIMERS**

Nucleic acid isolated or synthesized in accordance with the sequences described herein have utility as primers for the amplification of *M. catarrhalis* nucleic acid. These nucleic acids may also have utility as primers for the amplification of nucleic acids in other *Moraxella* species. With respect to polymerase chain reaction (PCR) techniques, nucleic acid sequences of ≥ 10-15 nucleotides of the invention contained in the Sequence Listing have utility in conjunction with suitable enzymes and reagents to create copies of *M*.

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catarrhalis nucleic acid. More preferably, the sequence will comprise twenty or more nucleotides to convey stability to the hybridization product formed between the primer and the intended target molecules. Binding conditions of primers greater than 100 nucleotides are more difficult to control to obtain specificity. High fidelity PCR can be used to ensure a faithful DNA copy prior to expression. In addition, amplified products can be checked by conventional sequencing methods.

The copies can be used in diagnostic assays to detect specific sequences, including genes from *M. catarrhalis* and/or other *Moraxella* species. The copies can also be incorporated into cloning and expression vectors to generate polypeptides corresponding to the nucleic acid synthesized by PCR, as is described in greater detail herein.

The nucleic acids of the present invention find use as templates for the recombinant production of *M. catarrhalis* -derived peptides or polypeptides

#### **ANTISENSE**

Nucleic acid or nucleic acid-hybridizing derivatives isolated or synthesized in accordance with the sequences described herein have utility as antisense agents to prevent the expression of *M. catarrhalis* genes. These sequences also have utility as antisense agents to prevent expression of genes of other *Moraxella* species.

In one embodiment, nucleic acid or derivatives corresponding to *M. catarrhalis* nucleic acids is loaded into a suitable carrier such as a liposome or bacteriophage for introduction into bacterial cells. For example, a nucleic acid having twenty or more nucleotides is capable of binding to bacteria nucleic acid or bacteria messenger RNA. Preferably, the antisense nucleic acid is comprised of 20 or more nucleotides to provide necessary stability of a hybridization product of non-naturally occurring nucleic acid and bacterial nucleic acid and/or bacterial messenger RNA. Nucleic acid having a sequence greater than 1000 nucleotides in length is difficult to synthesize but can be generated by recombinant DNA techniques. Methods for loading antisense nucleic acid in liposomes is known in the art as exemplified by U.S. Patent 4,241,046 issued December 23, 1980 to Papahadjopoulos et al.

The present invention encompasses isolated polypeptides and nucleic acids derived from *M. catarrhalis* that are useful as reagents for diagnosis of bacterial infection, components of effective anti-bacterial vaccines, and/or as targets for anti-bacterial drugs, including anti-*M. catarrhalis* drugs.

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#### EXPRESSION OF M. CATARRHALIS NUCLEIC ACIDS

Table 2, which is appended herewith and which forms part of the present specification, provides a list of open reading frames (ORFs) in both strands and a putative identification of the particular function of a polypeptide which is encoded by each ORF, based on the homology match (determined by the BLASTP2 algorithm) of the predicted polypeptide with known proteins encoded by ORFs in other organisms. An ORF is a region of nucleic acid which encodes a polypeptide. This region may represent a portion of a coding sequence or a total sequence and was determined from stop to stop codons. The first column contains a designation for the ORF ("ORF Name"). The second and third columns list the SEQ ID numbers for the nucleic acid ("NT ID") and amino acid ("AA ID") sequences corresponding to each ORF, respectively. The fourth and fifth columns list the length of the nucleic acid ORF ("NT Length") and the length of the amino acid ORF ("AA Length"), respectively. The nucleotide sequence corresponding to each ORF begins at the first nucleotide immediately following a stop codon and ends at the nucleotide immediately preceding the next downstream stop codon in the same reading frame. It will be recognized by one skilled in the art that the natural translation initiation sites will correspond to ATG, GTG, or TTG codons located within the ORFs. The natural initiation sites depend not only on the sequence of a start codon but also on the context of the DNA sequence adjacent to the start codon. Usually, a recognizable ribosome binding site is found within 20 nucleotides upstream from the initiation codon. In some cases where genes are translationally coupled and coordinately expressed together in "operons", ribosome binding sites are not present, but the initiation codon of a downstream gene may occur very close to, or overlap, the stop codon of the an upstream gene in the same operon. The correct start codons can be generally identified without undue experimentation because only a few codons need be tested. It is recognized that the translational machinery in bacteria initiates all polypeptide chains with

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the amino acid methionine, regardless of the sequence of the start codon. In some cases, polypeptides are post-translationally modified, resulting in an N-terminal amino acid other than methionine in vivo. The sixth and seventh columns provide metrics for assessing the likelihood of the homology match (determined by the BLASTP2 algorithm), as is known in the art, to the genes indicated in the description frame ("Description") defined further below. These genes in the Description were identified when the designated ORF was compared against a comprehensive non-redundant protein database. Specifically, the sixth column represents the Blast Score ("Score") for the match (a higher score is a better match), and the seventh column represents the probability ("Probability") for the match (the probability that such a match can have occurred by chance; the lower the value, the more likely the match is valid). If a BLASTP2 score of less than 100 was obtained, no value is reported in the table. The remaining fields below the columns contain additional information relating to the potential function of the sequence based on the BLASTP2 analysis. Where a match was discovered, the field "Protein name" list the protein's name identified from the match. In addition, one skilled in the art would be able to identify the match and elucidate its function using the "Locus name" and where available the accession number, "Acc#" from the database. Lastly, one skilled in the art would appreciate the "Description" field to further describe the potential function of the protein based on this analysis. This information allows one of ordinary skill in the art to determine a potential use for each identified coding sequence and, as a result, allows to use the polypeptides of the present invention for commercial and industrial purposes.

Using the information provided in SEQ ID NO: 1 - SEQ ID NO: 1920, SEQ ID NO: 1921 - SEQ ID NO: 3840 and in Table 2 together with routine cloning and sequencing methods, one of ordinary skill in the art will be able to clone and sequence all the nucleic acid fragments of interest including open reading frames (ORFs) encoding a large variety of proteins of *M. catarrhalis*.

Nucleic acid isolated or synthesized in accordance with the sequences described herein have utility to generate polypeptides. The nucleic acid of the invention exemplified in SEQ ID NO: 1 - SEQ ID NO: 1920 and in Table 2 or fragments of said nucleic acid encoding active portions of *M. catarrhalis* polypeptides can be cloned into suitable vectors

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or used to isolate nucleic acid. The isolated nucleic acid is combined with suitable DNA linkers and cloned into a suitable vector.

The function of a specific gene or operon can be ascertained by expression in a bacterial strain under conditions where the activity of the gene product(s) specified by the gene or operon in question can be specifically measured. Alternatively, a gene product may be produced in large quantities in an expressing strain for use as an antigen, an industrial reagent, for structural studies, etc. This expression can be accomplished in a mutant strain which lacks the activity of the gene to be tested, or in a strain that does not produce the same gene product(s). This includes, but is not limited to, Eucaryotic species such as the yeast *Saccharomyces cerevisiae, Methanobacterium* strains or other Archaea, and Eubacteria such as *E. coli*, *B. Subtilis*, *S. Aureus*, *S. Pneumonia* or *Pseudomonas putida*. In some cases the expression host will utilize the natural *M. catarrhalis* promoter whereas in others, it will be necessary to drive the gene with a promoter sequence derived from the expressing organism (e.g., an *E. coli* beta-galactosidase promoter for expression in *E. coli*).

To express a gene product using the natural *M. catarrhalis* promoter, a procedure such as the following can be used. A restriction fragment containing the gene of interest, together with its associated natural promoter element and regulatory sequences (identified using the DNA sequence data) is cloned into an appropriate recombinant plasmid containing an origin of replication that functions in the host organism and an appropriate selectable marker. This can be accomplished by a number of procedures known to those skilled in the art. It is most preferably done by cutting the plasmid and the fragment to be cloned with the same restriction enzyme to produce compatible ends that can be ligated to join the two pieces together. The recombinant plasmid is introduced into the host organism by, for example, electroporation and cells containing the recombinant plasmid are identified by selection for the marker on the plasmid. Expression of the desired gene product is detected using an assay specific for that gene product.

In the case of a gene that requires a different promoter, the body of the gene (coding sequence) is specifically excised and cloned into an appropriate expression plasmid. This subcloning can be done by several methods, but is most easily accomplished by PCR

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amplification of a specific fragment and ligation into an expression plasmid after treating the PCR product with a restriction enzyme or exonuclease to create suitable ends for cloning.

A suitable host cell for expression of a gene can be any procaryotic or eucaryotic cell. Suitable methods for transforming host cells can be found in Sambrook et al. (Molecular Cloning: A Laboratory Manual, 2nd Edition, Cold Spring Harbor Laboratory Press (1989)), and other laboratory textbooks.

For example, a host cell transfected with a nucleic acid vector directing expression of a nucleotide sequence encoding an *M. catarrhalis* polypeptide can be cultured under appropriate conditions to allow expression of the polypeptide to occur. Suitable media for cell culture are well known in the art. Polypeptides of the invention can be isolated from cell culture medium, host cells, or both using techniques known in the art for purifying proteins including ion-exchange chromatography, gel filtration chromatography, ultrafiltration, electrophoresis, and immunoaffinity purification with antibodies specific for such polypeptides. Additionally, in many situations, polypeptides can be produced by chemical cleavage of a native protein (e.g., tryptic digestion) and the cleavage products can then be purified by standard techniques.

In the case of membrane bound proteins, these can be isolated from a host cell by contacting a membrane-associated protein fraction with a detergent forming a solubilized complex, where the membrane-associated protein is no longer entirely embedded in the membrane fraction and is solubilized at least to an extent which allows it to be chromatographically isolated from the membrane fraction. Chromatographic techniques which can be used in the final purification step are known in the art and include hydrophobic interaction, lectin affinity, ion exchange, dye affinity and immunoaffinity.

One strategy to maximize recombinant *M. catarrhalis* peptide expression in *E. coli* is to express the protein in a host bacteria with an impaired capacity to proteolytically cleave the recombinant protein (Gottesman, S., Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, California (1990) 119-128). Another strategy would be to alter the nucleic acid encoding an *M. catarrhalis* peptide to be inserted into an expression vector so that the individual codons for each amino acid would be those preferentially utilized in highly expressed *E. coli* proteins (Wada et al., (1992) *Nuc. Acids* 

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Res. 20:2111-2118). Such alteration of nucleic acids of the invention can be carried out by standard DNA synthesis techniques.

The nucleic acids of the invention can also be chemically synthesized using standard techniques. Various methods of chemically synthesizing polydeoxynucleotides are known, including solid-phase synthesis which, like peptide synthesis, has been fully automated in commercially available DNA synthesizers (See, e.g., Itakura et al. U.S. Patent No. 4,598,049; Caruthers et al. U.S. Patent No. 4,458,066; and Itakura U.S. Patent Nos. 4,401,796 and 4,373,071, incorporated by reference herein).

The present invention provides a library of *M. catarrhalis* -derived nucleic acid sequences. The libraries provide probes, primers, and markers which can be used as markers in epidemiological studies. The present invention also provides a library of *M. catarrhalis* - derived nucleic acid sequences which comprise or encode targets for therapeutic drugs.

Nucleic acids comprising any of the sequences disclosed herein or sub-sequences thereof can be prepared by standard methods using the nucleic acid sequence information provided in SEQ ID NO: 1 - SEQ ID NO: 1920. For example, DNA can be chemically synthesized using, e.g., the phosphoramidite solid support method of Matteucci *et al.*, 1981, *J. Am. Chem. Soc.* 103:3185, the method of Yoo *et al.*, 1989, *J. Biol. Chem.* 764:17078, or other well known methods. This can be done by sequentially linking a series of oligonucleotide cassettes comprising pairs of synthetic oligonucleotides, as described below.

Of course, due to the degeneracy of the genetic code, many different nucleotide sequences can encode polypeptides having the amino acid sequences defined by SEQ ID NO: 1921 - SEQ ID NO: 3840 or sub-sequences thereof. The codons can be selected for optimal expression in prokaryotic or eukaryotic systems. Such degenerate variants are also encompassed by this invention.

Insertion of nucleic acids (typically DNAs) encoding the polypeptides of the invention into a vector is easily accomplished when the termini of both the DNAs and the vector comprise compatible restriction sites. If this cannot be done, it may be necessary to modify the termini of the DNAs and/or vector by digesting back single-stranded DNA overhangs generated by restriction endonuclease cleavage to produce blunt ends, or to

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achieve the same result by filling in the single-stranded termini with an appropriate DNA polymerase.

Alternatively, any site desired may be produced, e.g., by ligating nucleotide sequences (linkers) onto the termini. Such linkers may comprise specific oligonucleotide sequences that define desired restriction sites. Restriction sites can also be generated by the use of the polymerase chain reaction (PCR). See, e.g., Saiki et al., 1988, Science 239:48. The cleaved vector and the DNA fragments may also be modified if required by homopolymeric tailing.

The nucleic acids of the invention may be isolated directly from cells. Alternatively, the polymerase chain reaction (PCR) method can be used to produce the nucleic acids of the invention, using either chemically synthesized strands or genomic material as templates. Primers used for PCR can be synthesized using the sequence information provided herein and can further be designed to introduce appropriate new restriction sites, if desirable, to facilitate incorporation into a given vector for recombinant expression.

The nucleic acids of the present invention may be flanked by natural *M. catarrhalis* regulatory sequences, or may be associated with heterologous sequences, including promoters, enhancers, response elements, signal sequences, polyadenylation sequences, introns, 5'- and 3'- noncoding regions, and the like. The nucleic acids may also be modified by many means known in the art. Non-limiting examples of such modifications include methylation, "caps", substitution of one or more of the naturally occurring nucleotides with an analog, internucleotide modifications such as, for example, those with uncharged linkages (e.g., methyl phosphonates, phosphotriesters, phosphoroamidates, carbamates, etc.) and with charged linkages (e.g., phosphorothioates, phosphorodithioates, etc.). Nucleic acids may contain one or more additional covalently linked moieties, such as, for example, proteins (e.g., nucleases, toxins, antibodies, signal peptides, poly-L-lysine, etc.), intercalators (e.g., acridine, psoralen, etc.), chelators (e.g., metals, radioactive metals, iron, oxidative metals, etc.), and alkylators. PNAs are also included. The nucleic acid may be derivatized by formation of a methyl or ethyl phosphotriester or an alkyl phosphoramidate linkage. Furthermore, the nucleic acid sequences of the present invention may also be modified with

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a label capable of providing a detectable signal, either directly or indirectly. Exemplary labels include radioisotopes, fluorescent molecules, biotin, and the like.

The invention also provides nucleic acid vectors comprising the disclosed *M.* catarrhalis -derived sequences or derivatives or fragments thereof. A large number of vectors, including plasmid and bacterial vectors, have been described for replication and/or expression in a variety of eukaryotic and prokaryotic hosts, and may be used for cloning or protein expression.

The encoded *M. catarrhalis* polypeptides may be expressed by using many known vectors, such as pUC plasmids, pET plasmids (Novagen, Inc., Madison, WI), or pRSET or pREP (Invitrogen, San Diego, CA), and many appropriate host cells, using methods disclosed or cited herein or otherwise known to those skilled in the relevant art. The particular choice of vector/host is not critical to the practice of the invention.

Recombinant cloning vectors will often include one or more replication systems for cloning or expression, one or more markers for selection in the host, e.g. antibiotic resistance, and one or more expression cassettes. The inserted *M. catarrhalis* coding sequences may be synthesized by standard methods, isolated from natural sources, or prepared as hybrids, etc. Ligation of the *M. catarrhalis* coding sequences to transcriptional regulatory elements and/or to other amino acid coding sequences may be achieved by known methods. Suitable host cells may be transformed/transfected/infected as appropriate by any suitable method including electroporation, CaCl<sub>2</sub> mediated DNA uptake, bacterial infection, microprojectile, or other established methods.

Appropriate host cells include bacteria, archebacteria, fungi, especially yeast, and plant and animal cells, especially mammalian cells. Of particular interest are *M. catarrhalis*, *E. coli*, *B. Subtilis*, *Saccharomyces cerevisiae*, *Saccharomyces carlsbergensis*, *Schizosaccharomyces pombi*, *SF9* cells, C129 cells, 293 cells, *Neurospora*, and CHO cells, COS cells. He la cells, and immortalized mammalian musclaid and la cells and immortalized mammalian musclaid and la cells.

COS cells, HeLa cells, and immortalized mammalian myeloid and lymphoid cell lines. Preferred replication systems include M13, ColE1, SV40, baculovirus, lambda, adenovirus, and the like. A large number of transcription initiation and termination regulatory regions have been isolated and shown to be effective in the transcription and translation of

30 heterologous proteins in the various hosts. Examples of these regions, methods of isolation,

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manner of manipulation, etc. are known in the art. Under appropriate expression conditions, host cells can be used as a source of recombinantly produced *M. catarrhalis* -derived peptides and polypeptides.

Advantageously, vectors may also include a transcription regulatory element (i.e., a promoter) operably linked to the M. catarrhalis portion. The promoter may optionally contain operator portions and/or ribosome binding sites. Non-limiting examples of bacterial promoters compatible with E. coli include: b-lactamase (penicillinase) promoter; lactose promoter; tryptophan (trp) promoter; araBAD (arabinose) operon promoter; lambda-derived P<sub>I</sub> promoter and N gene ribosome binding site; and the hybrid tac promoter derived from sequences of the trp and lac UV5 promoters. Non-limiting examples of yeast promoters include 3-phosphoglycerate kinase promoter, glyceraldehyde-3-phosphate dehydrogenase (GAPDH) promoter, galactokinase (GAL1) promoter, galactoepimerase promoter, and alcohol dehydrogenase (ADH) promoter. Suitable promoters for mammalian cells include without limitation viral promoters such as that from Simian Virus 40 (SV40), Rous sarcoma virus (RSV), adenovirus (ADV), and bovine papilloma virus (BPV). Mammalian cells may also require terminator sequences, polyA addition sequences and enhancer sequences to increase expression. Sequences which cause amplification of the gene may also be desirable. Furthermore, sequences that facilitate secretion of the recombinant product from cells, including, but not limited to, bacteria, yeast, and animal cells, such as secretory signal sequences and/or prohormone pro region sequences, may also be included. These sequences are well described in the art.

Nucleic acids encoding wild-type or variant *M. catarrhalis* -derived polypeptides may also be introduced into cells by recombination events. For example, such a sequence can be introduced into a cell, and thereby effect homologous recombination at the site of an endogenous gene or a sequence with substantial identity to the gene. Other recombination-based methods such as nonhomologous recombinations or deletion of endogenous genes by homologous recombination may also be used.

The nucleic acids of the present invention find use as templates for the recombinant production of *M. catarrhalis* -derived peptides or polypeptides.

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## IDENTIFICATION AND USE OF M. CATARRHALIS NUCLEIC ACID. SEQUENCES

The disclosed *M. catarrhalis* polypeptide and nucleic acid sequences, or other sequences that are contained within ORFs, including complete protein-coding sequences, of which any of the disclosed *M. catarrhalis* -specific sequences forms a part, are useful as target components for diagnosis and/or treatment of *M. catarrhalis* - caused infection

It will be understood that the sequence of an entire protein-coding sequence of which each disclosed nucleic acid sequence forms a part can be isolated and identified based on each disclosed sequence. This can be achieved, for example, by using an isolated nucleic acid encoding the disclosed sequence, or fragments thereof, to prime a sequencing reaction with genomic *M. catarrhalis* DNA as template; this is followed by sequencing the amplified product. The isolated nucleic acid encoding the disclosed sequence, or fragments thereof, can also be hybridized to *M. catarrhalis* genomic libraries to identify clones containing additional complete segments of the protein-coding sequence of which the shorter sequence forms a part. Then, the entire protein-coding sequence, or fragments thereof, or nucleic acids encoding all or part of the sequence, or sequence-conservative or function-conservative variants thereof, may be employed in practicing the present invention.

Preferred sequences are those that are useful in diagnostic and/or therapeutic applications. Diagnostic applications include without limitation nucleic-acid-based and antibody-based methods for detecting bacterial infection. Therapeutic applications include without limitation vaccines, passive immunotherapy, and drug treatments directed against gene products that are both unique to bacteria and essential for growth and/or replication of bacteria.

# IDENTIFICATION OF NUCLEIC ACIDS ENCODING VACCINE COMPONENTS AND TARGETS FOR AGENTS EFFECTIVE AGAINST M. CATARRHALIS

The disclosed *M. catarrhalis* genome sequence includes segments that direct the synthesis of ribonucleic acids and polypeptides, as well as origins of replication, promoters, other types of regulatory sequences, and intergenic nucleic acids. The invention encompasses nucleic acids encoding immunogenic components of vaccines and targets for agents effective against *M. catarrhalis*. Identification of said immunogenic components

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involved in the determination of the function of the disclosed sequences, which can be achieved using a variety of approaches. Non-limiting examples of these approaches are described briefly below.

### 5 HOMOLOGY TO KNOWN SEQUENCES:

Computer-assisted comparison of the disclosed M. catarrhalis sequences with previously reported sequences present in publicly available databases is useful for identifying functional M. catarrhalis nucleic acid and polypeptide sequences. It will be understood that protein-coding sequences, for example, may be compared as a whole, and that a high degree of sequence homology between two proteins (such as, for example, >80-90%) at the amino acid level indicates that the two proteins also possess some degree of functional homology, such as, for example, among enzymes involved in metabolism, DNA synthesis, or cell wall synthesis, and proteins involved in transport, cell division, etc. In addition, many structural features of particular protein classes have been identified and correlate with specific consensus sequences, such as, for example, binding domains for nucleotides, DNA, metal ions, and other small molecules; sites for covalent modifications such as phosphorylation, acylation, and the like; sites of protein:protein interactions, etc. These consensus sequences may be quite short and thus may represent only a fraction of the entire protein-coding sequence. Identification of such a feature in an M. catarrhalis sequence is therefore useful in determining the function of the encoded protein and identifying useful targets of antibacterial drugs.

Of particular relevance to the present invention are structural features that are common to secretory, transmembrane, and surface proteins, including secretion signal peptides and hydrophobic transmembrane domains. *M. catarrhalis* proteins identified as containing putative signal sequences and/or transmembrane domains are useful as immunogenic components of vaccines.

Targets for therapeutic drugs according to the invention include, but are not limited to, polypeptides of the invention, whether unique to *M. catarrhalis* or not, that are essential for growth and/or viability of *M. catarrhalis* under at least one growth condition.

30 Polypeptides essential for growth and/or viability can be determined by examining the effect

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of deleting and/or disrupting the genes, i.e., by so-called gene "knockout". Alternatively, genetic footprinting can be used (Smith et al., 1995, Proc. Natl. Acad. Sci. USA 92:5479-6433; Published International Application WO 94/26933; U.S. Patent No. 5,612,180). Still other methods for assessing essentiality includes the ability to isolate conditional lethal mutations in the specific gene (e.g., temperature sensitive mutations). Other useful targets for therapeutic drugs, which include polypeptides that are not essential for growth or viability per se but lead to loss of viability of the cell, can be used to target therapeutic agents to cells.

### 10 STRAIN-SPECIFIC SEQUENCES:

Because of the evolutionary relationship between different *M. catarrhalis* strains, it is believed that the presently disclosed *M. catarrhalis* sequences are useful for identifying, and/or discriminating between, previously known and new *M. catarrhalis* strains. It is believed that other *M. catarrhalis* strains will exhibit at least about 70% sequence homology with the presently disclosed sequence. Systematic and routine analyses of DNA sequences derived from samples containing *M. catarrhalis* strains, and comparison with the present sequence allows for the identification of sequences that can be used to discriminate between strains, as well as those that are common to all *M. catarrhalis* strains. In one embodiment, the invention provides nucleic acids, including probes, and peptide and polypeptide sequences that discriminate between different strains of *M. catarrhalis*. Strain-specific components can also be identified functionally by their ability to elicit or react with antibodies that selectively recognize one or more *M. catarrhalis* strains.

In another embodiment, the invention provides nucleic acids, including probes, and peptide and polypeptide sequences that are common to all *M. catarrhalis* strains but are not found in other bacterial species.

#### M. CATARRHALIS POLYPEPTIDES

This invention encompasses isolated *M. catarrhalis* polypeptides encoded by the disclosed *M. catarrhalis* genomic sequences, including the polypeptides of the invention contained in the Sequence Listing. Polypeptides of the invention are preferably at least

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about 5 amino acid residues in length. Using the DNA sequence information provided herein, the amino acid sequences of the polypeptides encompassed by the invention can be deduced using methods well-known in the art. It will be understood that the sequence of an entire nucleic acid encoding an *M. catarrhalis* polypeptide can be isolated and identified based on an ORF that encodes only a fragment of the cognate protein-coding region. This can be achieved, for example, by using the isolated nucleic acid encoding the ORF, or fragments thereof, to prime a polymerase chain reaction with genomic *M. catarrhalis* DNA as template; this is followed by sequencing the amplified product.

The polypeptides of the present invention, including function-conservative variants of the disclosed ORFs, may be isolated from wild-type or mutant *M. catarrhalis* cells, or from heterologous organisms or cells (including, but not limited to, bacteria, fungi, insect, plant, and mammalian cells) including *M. catarrhalis* into which an *M. catarrhalis* -derived protein-coding sequence has been introduced and expressed. Furthermore, the polypeptides may be part of recombinant fusion proteins.

M. catarrhalis polypeptides of the invention can be chemically synthesized using commercially automated procedures such as those referenced herein, including, without limitation, exclusive solid phase synthesis, partial solid phase methods, fragment condensation or classical solution synthesis. The polypeptides are preferably prepared by solid phase peptide synthesis as described by Merrifield, 1963, J. Am. Chem. Soc. 85:2149. The synthesis is carried out with amino acids that are protected at the alpha-amino terminus. Trifunctional amino acids with labile side-chains are also protected with suitable groups to prevent undesired chemical reactions from occurring during the assembly of the polypeptides. The alpha-amino protecting group is selectively removed to allow subsequent reaction to take place at the amino-terminus. The conditions for the removal of the alpha-amino protecting group do not remove the side-chain protecting groups.

Methods for polypeptide purification are well-known in the art, including, without limitation, preparative disc-gel electrophoresis, isoelectric focusing, HPLC, reversed-phase HPLC, gel filtration, ion exchange and partition chromatography, and countercurrent distribution. For some purposes, it is preferable to produce the polypeptide in-a recombinant system in which the *M. catarrhalis* protein contains an additional sequence tag that

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facilitates purification, such as, but not limited to, a polyhistidine sequence. The polypeptide can then be purified from a crude lysate of the host cell by chromatography on an appropriate solid-phase matrix. Alternatively, antibodies produced against an *M. catarrhalis* protein or against peptides derived therefrom can be used as purification reagents. Other purification methods are possible.

The present invention also encompasses derivatives and homologues of *M.* catarrhalis -encoded polypeptides. For some purposes, nucleic acid sequences encoding the peptides may be altered by substitutions, additions, or deletions that provide for functionally equivalent molecules, i.e., function-conservative variants. For example, one or more amino acid residues within the sequence can be substituted by another amino acid of similar properties, such as, for example, positively charged amino acids (arginine, lysine, and histidine); negatively charged amino acids (aspartate and glutamate); polar neutral amino acids; and non-polar amino acids.

The isolated polypeptides may be modified by, for example, phosphorylation, sulfation, acylation, or other protein modifications. They may also be modified with a label capable of providing a detectable signal, either directly or indirectly, including, but not limited to, radioisotopes and fluorescent compounds.

To identify *M. catarrhalis* -derived polypeptides for use in the present invention, essentially the complete genomic sequence of a virulent, methicillin-resistant isolate of *M. catarrhalis* isolate was analyzed. While, in very rare instances, a nucleic acid sequencing error may be revealed, resolving a rare sequencing error is well within the art, and such an occurrence will not prevent one skilled in the art from practicing the invention.

Also encompassed are any *M. catarrhalis* polypeptide sequences that are contained within the open reading frames (ORFs), including complete protein-coding sequences, of which any of SEQ ID NO: 1 - SEQ ID NO: 1920 forms a part. Table 2, which is appended herewith and which forms part of the present specification, provides a putative identification of the particular function of a polypeptide which is encoded by each ORF, based on the homology match (determined by the BLAST algorithm) of the predicted polypeptide with known proteins encoded by ORFs in other organisms. As a result, one skilled in the art can

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use the polypeptides of the present invention for commercial and industrial purposes consistent with the type of putative identification of the polypeptide.

The present invention provides a library of *M. catarrhalis* -derived polypeptide sequences, and a corresponding library of nucleic acid sequences encoding the polypeptides, wherein the polypeptides themselves, or polypeptides contained within ORFs of which they form a part, comprise sequences that are contemplated for use as components of vaccines. Non-limiting examples of such sequences are listed by SEQ ID NO in Table 2, which is appended herewith and which forms part of the present specification.

The present invention also provides a library of *M. catarrhalis* -derived polypeptide sequences, and a corresponding library of nucleic acid sequences encoding the polypeptides, wherein the polypeptides themselves, or polypeptides contained within ORFs of which they form a part, comprise sequences lacking homology to any known prokaryotic or eukaryotic sequences. Such libraries provide probes, primers, and markers which can be used to diagnose *M. catarrhalis* infection, including use as markers in epidemiological studies. Non-limiting examples of such sequences are listed by SEQ ID NO in Table 2, which is appended hereto and part hereof.

The present invention also provides a library of *M. catarrhalis* -derived polypeptide sequences, and a corresponding library of nucleic acid sequences encoding the polypeptides, wherein the polypeptides themselves, or polypeptides contained within ORFs of which they form a part, comprise targets for therapeutic drugs.

## SPECIFIC EXAMPLE: DETERMINATION OF *MORAXELLA* PROTEIN ANTIGENS FOR ANTIBODY AND VACCINE DEVELOPMENT

The selection of Moraxella protein antigens for vaccine development can be derived from the nucleic acids encoding *M. catarrhalis* polypeptides. First, the ORF's can be analyzed for homology to other known exported or membrane proteins and analyzed using the discriminant analysis described by Klein, et al. (Klein, P., Kanehsia, M., and DeLisi, C. (1985) *Biochimica et Biophysica Acta* 815, 468-476) for predicting exported and membrane proteins.

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Homology searches can be performed using the BLAST algorithm contained in the Wisconsin Sequence Analysis Package (Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711) to compare each predicted ORF amino acid sequence with all sequences found in the current GenBank, SWISS-PROT and PIR databases. BLAST searches for local alignments between the ORF and the databank sequences and reports a probability score which indicates the probability of finding this sequence by chance in the database. ORF's with significant homology (e.g. probabilities lower than  $1\times10^{-6}$  that the homology is only due to random chance) to membrane or exported proteins represent protein antigens for vaccine development. Possible functions can be provided to *M. catarrhalis* genes based on sequence homology to genes cloned in other organisms.

Discriminant analysis (Klein, et al. supra) can be used to examine the ORF amino acid sequences. This algorithm uses the intrinsic information contained in the ORF amino acid sequence and compares it to information derived from the properties of known membrane and exported proteins. This comparison predicts which proteins will be exported, membrane associated or cytoplasmic. ORF amino acid sequences identified as exported or membrane associated by this algorithm are likely protein antigens for vaccine development.

## PRODUCTION OF FRAGMENTS AND ANALOGS OF *M. CATARRHALIS* NUCLEIC ACIDS AND POLYPEPTIDES

Based on the discovery of the *M. catarrhalis* gene products of the invention provided in the Sequence Listing, one skilled in the art can alter the disclosed structure of *M. catarrhalis* genes, e.g., by producing fragments or analogs, and test the newly produced structures for activity. Examples of techniques known to those skilled in the relevant art which allow the production and testing of fragments and analogs are discussed below. These, or analogous methods can be used to make and screen libraries of polypeptides, e.g., libraries of random peptides or libraries of fragments or analogs of cellular proteins for the ability to bind *M. catarrhalis* polypeptides. Such screens are useful for the identification of inhibitors of *M. catarrhalis*.

### GENERATION OF FRAGMENTS

Fragments of a protein can be produced in several ways, e.g., recombinantly, by proteolytic digestion, or by chemical synthesis. Internal or terminal fragments of a polypeptide can be generated by removing one or more nucleotides from one end (for a terminal fragment) or both ends (for an internal fragment) of a nucleic acid which encodes the polypeptide. Expression of the mutagenized DNA produces polypeptide fragments. Digestion with "end-nibbling" endonucleases can thus generate DNAs which encode an array of fragments. DNAs which encode fragments of a protein can also be generated by random shearing, restriction digestion or a combination of the above-discussed methods.

Fragments can also be chemically synthesized using techniques known in the art such as conventional Merrifield solid phase f-Moc or t-Boc chemistry. For example, peptides of the present invention may be arbitrarily divided into fragments of desired length with no overlap of the fragments, or divided into overlapping fragments of a desired length.

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### ALTERATION OF NUCLEIC ACIDS AND POLYPEPTIDES: RANDOM METHODS

Amino acid sequence variants of a protein can be prepared by random mutagenesis of DNA which encodes a protein or a particular domain or region of a protein. Useful methods include PCR mutagenesis and saturation mutagenesis. A library of random amino acid sequence variants can also be generated by the synthesis of a set of degenerate oligonucleotide sequences. (Methods for screening proteins in a library of variants are elsewhere herein).

### PCR MUTAGENESIS

In PCR mutagenesis, reduced Taq polymerase fidelity is used to introduce random mutations into a cloned fragment of DNA (Leung et al., 1989, *Technique* 1:11-15). The DNA region to be mutagenized is amplified using the polymerase chain reaction (PCR) under conditions that reduce the fidelity of DNA synthesis by Taq DNA polymerase, e.g., by using a dGTP/dATP ratio of five and adding Mn 2+ to the PCR reaction. The pool of

amplified DNA fragments are inserted into appropriate cloning vectors to provide random mutant libraries.

#### SATURATION MUTAGENESIS

Saturation mutagenesis allows for the rapid introduction of a large number of single base substitutions into cloned DNA fragments (Mayers et al., 1985, Science 229:242). This technique includes generation of mutations, e.g., by chemical treatment or irradiation of single-stranded DNA in vitro, and synthesis of a complimentary DNA strand. The mutation frequency can be modulated by modulating the severity of the treatment, and essentially all possible base substitutions can be obtained. Because this procedure does not involve a genetic selection for mutant fragments both neutral substitutions, as well as those that alter function, are obtained. The distribution of point mutations is not biased toward conserved sequence elements.

### 15 DEGENERATE OLIGONUCLEOTIDES

A library of homologs can also be generated from a set of degenerate oligonucleotide sequences. Chemical synthesis of a degenerate sequences can be carried out in an automatic DNA synthesizer, and the synthetic genes then ligated into an appropriate expression vector. The synthesis of degenerate oligonucleotides is known in the art (see for example, Narang, SA (1983) Tetrahedron 39:3; Itakura et al. (1981) Recombinant DNA, Proc 3rd Cleveland Sympos. Macromolecules, ed. AG Walton, Amsterdam: Elsevier pp273-289; Itakura et al. (1984) Annu. Rev. Biochem. 53:323; Itakura et al. (1984) Science 198:1056; Ike et al. (1983) Nucleic Acid Res. 11:477. Such techniques have been employed in the directed evolution of other proteins (see, for example, Scott et al. (1990) Science 249:386-390; Roberts et al. (1992) PNAS 89:2429-2433; Devlin et al. (1990) Science 249: 404-406; Cwirla et al. (1990)

PNAS 87: 6378-6382; as well as U.S. Patents Nos. 5,223,409, 5,198,346, and 5,096,815).

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## ALTERATION OF NUCLEIC ACIDS AND POLYPEPTIDES: METHODS FOR DIRECTED MUTAGENESIS

Non-random or directed, mutagenesis techniques can be used to provide specific sequences or mutations in specific regions. These techniques can be used to create variants which include, e.g., deletions, insertions, or substitutions, of residues of the known amino acid sequence of a protein. The sites for mutation can be modified individually or in series, e.g., by (1) substituting first with conserved amino acids and then with more radical choices depending upon results achieved, (2) deleting the target residue, or (3) inserting residues of the same or a different class adjacent to the located site, or combinations of options 1-3.

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### ALANINE SCANNING MUTAGENESIS

Alanine scanning mutagenesis is a useful method for identification of certain residues or regions of the desired protein that are preferred locations or domains for mutagenesis, Cunningham and Wells (*Science* 244:1081-1085, 1989). In alanine scanning, a residue or group of target residues are identified (e.g., charged residues such as Arg, Asp, His, Lys, and Glu) and replaced by a neutral or negatively charged amino acid (most preferably alanine or polyalanine). Replacement of an amino acid can affect the interaction of the amino acids with the surrounding aqueous environment in or outside the cell. Those domains demonstrating functional sensitivity to the substitutions are then refined by introducing further or other variants at or for the sites of substitution. Thus, while the site for introducing an amino acid sequence variation is predetermined, the nature of the mutation per se need not be predetermined. For example, to optimize the performance of a mutation at a given site, alanine scanning or random mutagenesis may be conducted at the target codon or region and the expressed desired protein subunit variants are screened for the optimal combination of desired activity.

### OLIGONUCLEOTIDE-MEDIATED MUTAGENESIS

Oligonucleotide-mediated mutagenesis is a useful method for preparing substitution, deletion, and insertion variants of DNA, see, e.g., Adelman et al., (DNA 2:183, 1983). Briefly, the desired DNA is altered by hybridizing an oligonucleotide encoding a mutation to

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a DNA template, where the template is the single-stranded form of a plasmid or bacteriophage containing the unaltered or native DNA sequence of the desired protein. After hybridization, a DNA polymerase is used to synthesize an entire second complementary strand of the template that will thus incorporate the oligonucleotide primer, and will code for the selected alteration in the desired protein DNA. Generally, oligonucleotides of at least about 25 nucleotides in length are used. An optimal oligonucleotide will have 12 to 15 nucleotides that are completely complementary to the template on either side of the nucleotide(s) coding for the mutation. This ensures that the oligonucleotide will hybridize properly to the single-stranded DNA template molecule. The oligonucleotides are readily synthesized using techniques known in the art such as that described by Crea et al. (Proc. Natl. Acad. Sci. USA, 75: 5765[1978]).

### CASSETTE MUTAGENESIS

Another method for preparing variants, cassette mutagenesis, is based on the technique described by Wells et al. (Gene, 34:315[1985]). The starting material is a plasmid 15 (or other vector) which includes the protein subunit DNA to be mutated. The codon(s) in the protein subunit DNA to be mutated are identified. There must be a unique restriction endonuclease site on each side of the identified mutation site(s). If no such restriction sites exist, they may be generated using the above-described oligonucleotide-mediated mutagenesis method to introduce them at appropriate locations in the desired protein subunit 20 DNA. After the restriction sites have been introduced into the plasmid, the plasmid is cut at these sites to linearize it. A double-stranded oligonucleotide encoding the sequence of the DNA between the restriction sites but containing the desired mutation(s) is synthesized using standard procedures. The two strands are synthesized separately and then hybridized together using standard techniques. This double-stranded oligonucleotide is referred to as the cassette. This cassette is designed to have 3' and 5' ends that are comparable with the ends of the linearized plasmid, such that it can be directly ligated to the plasmid. This plasmid now contains the mutated desired protein subunit DNA sequence.

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### COMBINATORIAL MUTAGENESIS

Combinatorial mutagenesis can also be used to generate mutants (Ladner et al., WO 88/06630). In this method, the amino acid sequences for a group of homologs or other related proteins are aligned, preferably to promote the highest homology possible. All of the amino acids which appear at a given position of the aligned sequences can be selected to create a degenerate set of combinatorial sequences. The variegated library of variants is generated by combinatorial mutagenesis at the nucleic acid level, and is encoded by a variegated gene library. For example, a mixture of synthetic oligonucleotides can be enzymatically ligated into gene sequences such that the degenerate set of potential sequences are expressible as individual peptides, or alternatively, as a set of larger fusion proteins containing the set of degenerate sequences.

## OTHER MODIFICATIONS OF M. CATARRHALIS NUCLEIC ACIDS AND POLYPEPTIDES

It is possible to modify the structure of an *M. catarrhalis* polypeptide for such purposes as increasing solubility, enhancing stability (e.g., shelf life *ex vivo* and resistance to proteolytic degradation *in vivo*). A modified *M. catarrhalis* protein or peptide can be produced in which the amino acid sequence has been altered, such as by amino acid substitution, deletion, or addition as described herein.

An *M. catarrhalis* peptide can also be modified by substitution of cysteine residues preferably with alanine, serine, threonine, leucine or glutamic acid residues to minimize dimerization via disulfide linkages. In addition, amino acid side chains of fragments of the protein of the invention can be chemically modified. Another modification is cyclization of the peptide.

In order to enhance stability and/or reactivity, an *M. catarrhalis* polypeptide can be modified to incorporate one or more polymorphisms in the amino acid sequence of the protein resulting from any natural allelic variation. Additionally, D-amino acids, non-natural amino acids, or non-amino acid analogs can be substituted or added to produce a modified protein within the scope of this invention. Furthermore, an *M. catarrhalis* polypeptide can be modified using polyethylene glycol (PEG) according to the method of A. Sehon and co-

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workers (Wie et al., supra) to produce a protein conjugated with PEG. In addition, PEG can be added during chemical synthesis of the protein. Other modifications of *M. catarrhalis* proteins include reduction/alkylation (Tarr, *Methods of Protein Microcharacterization*, J. E. Silver ed., Humana Press, Clifton NJ 155-194 (1986)); acylation (Tarr, supra); chemical coupling to an appropriate carrier (Mishell and Shiigi, eds, *Selected Methods in Cellular Immunology*, WH Freeman, San Francisco, CA (1980), U.S. Patent 4,939,239; or mild formalin treatment (Marsh, (1971) *Int. Arch. of Allergy and Appl. Immunol.*, 41: 199 - 215).

To facilitate purification and potentially increase solubility of an *M. catarrhalis* protein or peptide, it is possible to add an amino acid fusion moiety to the peptide backbone. For example, hexa-histidine can be added to the protein for purification by immobilized metal ion affinity chromatography (Hochuli, E. et al., (1988) *Bio/Technology*, 6: 1321 - 1325). In addition, to facilitate isolation of peptides free of irrelevant sequences, specific endoprotease cleavage sites can be introduced between the sequences of the fusion moiety and the peptide.

To potentially aid proper antigen processing of epitopes within an *M. catarrhalis* polypeptide, canonical protease sensitive sites can be engineered between regions, each comprising at least one epitope via recombinant or synthetic methods. For example, charged amino acid pairs, such as KK or RR, can be introduced between regions within a protein or fragment during recombinant construction thereof. The resulting peptide can be rendered sensitive to cleavage by cathepsin and/or other trypsin-like enzymes which would generate portions of the protein containing one or more epitopes. In addition, such charged amino acid residues can result in an increase in the solubility of the peptide.

### PRIMARY METHODS FOR SCREENING POLYPEPTIDES AND ANALOGS

Various techniques are known in the art for screening generated mutant gene products. Techniques for screening large gene libraries often include cloning the gene library into replicable expression vectors, transforming appropriate cells with the resulting library of vectors, and expressing the genes under conditions in which detection of a desired activity, e.g., in this case, binding to *M. catarrhalis* polypeptide or an interacting protein, facilitates relatively easy isolation of the vector encoding the gene whose product was

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detected. Each of the techniques described below is amenable to high through-put analysis for screening large numbers of sequences created, e.g., by random mutagenesis techniques.

#### TWO HYBRID SYSTEMS

Two hybrid assays such as the system described below (as with the other screening methods described herein), can be used to identify polypeptides, e.g., fragments or analogs of a naturally-occurring *M. catarrhalis* polypeptide, e.g., of cellular proteins, or of randomly generated polypeptides which bind to an *M. catarrhalis* protein. (The *M. catarrhalis* domain is used as the bait protein and the library of variants are expressed as prey fusion proteins.) In an analogous fashion, a two hybrid assay (as with the other screening methods described herein), can be used to find polypeptides which bind an *M. catarrhalis* polypeptide.

#### DISPLAY LIBRARIES

In one approach to screening assays, the Moraxella peptides are displayed on the surface of a cell or viral particle, and the ability of particular cells or viral particles to bind an appropriate receptor protein via the displayed product is detected in a "panning assay". For example, the gene library can be cloned into the gene for a surface membrane protein of a bacterial cell, and the resulting fusion protein detected by panning (Ladner et al., WO 88/06630; Fuchs et al. (1991) *Bio/Technology* 9:1370-1371; and Goward et al. (1992) *TIBS* 18:136-140). In a similar fashion, a detectably labeled ligand can be used to score for potentially functional peptide homologs. Fluorescently labeled ligands, e.g., receptors, can be used to detect homologs which retain ligand-binding activity. The use of fluorescently labeled ligands, allows cells to be visually inspected and separated under a fluorescence microscope, or, where the morphology of the cell permits, to be separated by a fluorescence-activated cell sorter.

A gene library can be expressed as a fusion protein on the surface of a viral particle. For instance, in the filamentous phage system, foreign peptide sequences can be expressed on the surface of infectious phage, thereby conferring two significant benefits. First, since these phage can be applied to affinity matrices at concentrations well over 10<sup>13</sup> phage per

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milliliter, a large number of phage can be screened at one time. Second, since each infectious phage displays a gene product on its surface, if a particular phage is recovered from an affinity matrix in low yield, the phage can be amplified by another round of infection. The group of almost identical *E. coli* filamentous phages, M13, fd., and f1, are most often used in phage display libraries. Either of the phage gIII or gVIII coat proteins can be used to generate fusion proteins without disrupting the ultimate packaging of the viral particle. Foreign epitopes can be expressed at the NH<sub>2</sub>-terminal end of pIII and phage bearing such epitopes recovered from a large excess of phage lacking this epitope (Ladner et al. PCT publication WO 90/02909; Garrard et al., PCT publication WO 92/09690; Marks et al. (1992) *J. Biol. Chem.* 267:16007-16010; Griffiths et al. (1993) *EMBO J* 12:725-734; Clackson et al. (1991) *Nature* 352:624-628; and Barbas et al. (1992) *PNAS* 89:4457-4461).

A common approach uses the maltose receptor of E. coli (the outer membrane protein, LamB) as a peptide fusion partner (Charbit et al. (1986) EMBO 5, 3029-3037). Oligonucleotides have been inserted into plasmids encoding the LamB gene to produce peptides fused into one of the extracellular loops of the protein. These peptides are available for binding to ligands, e.g., to antibodies, and can elicit an immune response when the cells are administered to animals. Other cell surface proteins, e.g., OmpA (Schorr et al. (1991) Vaccines 91, pp. 387-392), PhoE (Agterberg, et al. (1990) Gene 88, 37-45), and PAL (Fuchs et al. (1991) Bio/Tech 9, 1369-1372), as well as large bacterial surface structures have served as vehicles for peptide display. Peptides can be fused to pilin, a protein which polymerizes to form the pilus-a conduit for interbacterial exchange of genetic information (Thiry et al. (1989) Appl. Environ. Microbiol. 55, 984-993). Because of its role in interacting with other cells, the pilus provides a useful support for the presentation of peptides to the extracellular environment. Another large surface structure used for peptide display is the bacterial motive organ, the flagellum. Fusion of peptides to the subunit protein flagellin offers a dense array of many peptide copies on the host cells (Kuwajima et al. (1988) Bio/Tech. 6, 1080-1083). Surface proteins of other bacterial species have also served as peptide fusion partners. Examples include the Moraxella protein A and the outer membrane IgA protease of Neisseria (Hansson et al. (1992) J. Bacteriol. 174, 4239-4245 and Klauser et al. (1990) EMBO J. 9, 1991-1999).

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In the filamentous phage systems and the LamB system described above, the physical link between the peptide and its encoding DNA occurs by the containment of the DNA within a particle (cell or phage) that carries the peptide on its surface. Capturing the peptide captures the particle and the DNA within. An alternative scheme uses the DNA-binding protein LacI to form a link between peptide and DNA (Cull et al. (1992) PNAS USA 89:1865-1869). This system uses a plasmid containing the LacI gene with an oligonucleotide cloning site at its 3'-end. Under the controlled induction by arabinose, a LacI-peptide fusion protein is produced. This fusion retains the natural ability of LacI to bind to a short DNA sequence known as LacO operator (LacO). By installing two copies of LacO on the expression plasmid, the LacI-peptide fusion binds tightly to the plasmid that encoded it. Because the plasmids in each cell contain only a single oligonucleotide sequence and each cell expresses only a single peptide sequence, the peptides become specifically and stablely associated with the DNA sequence that directed its synthesis. The cells of the library are gently lysed and the peptide-DNA complexes are exposed to a matrix of immobilized receptor to recover the complexes containing active peptides. The associated plasmid DNA is then reintroduced into cells for amplification and DNA sequencing to determine the identity of the peptide ligands. As a demonstration of the practical utility of the method, a large random library of dodecapeptides was made and selected on a monoclonal antibody raised against the opioid peptide dynorphin B. A cohort of peptides was recovered, all related by a consensus sequence corresponding to a six-residue portion of dynorphin B. (Cull et al. (1992) Proc. Natl. Acad. Sci. U.S.A. 89-1869)

This scheme, sometimes referred to as peptides-on-plasmids, differs in two important ways from the phage display methods. First, the peptides are attached to the C-terminus of the fusion protein, resulting in the display of the library members as peptides having free carboxy termini. Both of the filamentous phage coat proteins, pIII and pVIII, are anchored to the phage through their C-termini, and the guest peptides are placed into the outward-extending N-terminal domains. In some designs, the phage-displayed peptides are presented right at the amino terminus of the fusion protein. (Cwirla, et al. (1990) *Proc. Natl. Acad. Sci. U.S.* A. 87, 6378-6382) A second difference is the set of biological biases affecting the population of peptides actually present in the libraries. The LacI fusion molecules are

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confined to the cytoplasm of the host cells. The phage coat fusions are exposed briefly to the cytoplasm during translation but are rapidly secreted through the inner membrane into the periplasmic compartment, remaining anchored in the membrane by their C-terminal hydrophobic domains, with the N-termini, containing the peptides, protruding into the periplasm while awaiting assembly into phage particles. The peptides in the LacI and phage libraries may differ significantly as a result of their exposure to different proteolytic activities. The phage coat proteins require transport across the inner membrane and signal peptidase processing as a prelude to incorporation into phage. Certain peptides exert a deleterious effect on these processes and are underrepresented in the libraries (Gallop et al. (1994) *J. Med. Chem.* 37(9):1233-1251). These particular biases are not a factor in the LacI display system.

The number of small peptides available in recombinant random libraries is enormous. Libraries of  $10^7$ - $10^9$  independent clones are routinely prepared. Libraries as large as  $10^{11}$  recombinants have been created, but this size approaches the practical limit for clone libraries. This limitation in library size occurs at the step of transforming the DNA containing randomized segments into the host bacterial cells. To circumvent this limitation, an *in vitro* system based on the display of nascent peptides in polysome complexes has recently been developed. This display library method has the potential of producing libraries 3-6 orders of magnitude larger than the currently available phage/phagemid or plasmid libraries. Furthermore, the construction of the libraries, expression of the peptides, and screening, is done in an entirely cell-free format.

In one application of this method (Gallop et al. (1994) *J. Med. Chem.* 37(9):1233-1251), a molecular DNA library encoding 10<sup>12</sup> decapeptides was constructed and the library expressed in an *E. coli* S30 *in vitro* coupled transcription/translation system. Conditions were chosen to stall the ribosomes on the mRNA, causing the accumulation of a substantial proportion of the RNA in polysomes and yielding complexes containing nascent peptides still linked to their encoding RNA. The polysomes are sufficiently robust to be affinity purified on immobilized receptors in much the same way as the more conventional recombinant peptide display libraries are screened. RNA from the bound complexes is

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recovered, converted to cDNA, and amplified by PCR to produce a template for the next round of synthesis and screening. The polysome display method can be coupled to the phage display system. Following several rounds of screening, cDNA from the enriched pool of polysomes was cloned into a phagemid vector. This vector serves as both a peptide expression vector, displaying peptides fused to the coat proteins, and as a DNA sequencing vector for peptide identification. By expressing the polysome-derived peptides on phage, one can either continue the affinity selection procedure in this format or assay the peptides on individual clones for binding activity in a phage ELISA, or for binding specificity in a completion phage ELISA (Barret, et al. (1992) *Anal. Biochem* 204,357-364). To identify the sequences of the active peptides one sequences the DNA produced by the phagemid host.

### SECONDARY SCREENING OF POLYPEPTIDES AND ANALOGS

The high through-put assays described above can be followed by secondary screens in order to identify further biological activities which will, e.g., allow one skilled in the art to differentiate agonists from antagonists. The type of a secondary screen used will depend on the desired activity that needs to be tested. For example, an assay can be developed in which the ability to inhibit an interaction between a protein of interest and its respective ligand can be used to identify antagonists from a group of peptide fragments isolated though one of the primary screens described above.

Therefore, methods for generating fragments and analogs and testing them for activity are known in the art. Once the core sequence of interest is identified, it is routine for one skilled in the art to obtain analogs and fragments.

### PEPTIDE MIMETICS OF M. CATARRHALIS POLYPEPTIDES

The invention also provides for reduction of the protein binding domains of the subject *M. catarrhalis* polypeptides to generate mimetics, e.g. peptide or non-peptide agents. The peptide mimetics are able to disrupt binding of a polypeptide to its counter ligand, e.g., in the case of an *M. catarrhalis* polypeptide binding to a naturally occurring ligand. The critical residues of a subject *M. catarrhalis* polypeptide which are involved in molecular recognition of a polypeptide can be determined and used to generate *M. catarrhalis* -derived

peptidomimetics which competitively or noncompetitively inhibit binding of the *M.* catarrhalis polypeptide with an interacting polypeptide (see, for example, European patent applications EP-412,762A and EP-B31,080A).

For example, scanning mutagenesis can be used to map the amino acid residues of a particular M. catarrhalis polypeptide involved in binding an interacting polypeptide, 5 peptidomimetic compounds (e.g. diazepine or isoquinoline derivatives) can be generated which mimic those residues in binding to an interacting polypeptide, and which therefore can inhibit binding of an M. catarrhalis polypeptide to an interacting polypeptide and thereby interfere with the function of M. catarrhalis polypeptide. For instance, non-hydrolyzable 10 peptide analogs of such residues can be generated using benzodiazepine (e.g., see Freidinger et al. in Peptides: Chemistry and Biology, G.R. Marshall ed., ESCOM Publisher: Leiden, Netherlands, 1988), azepine (e.g., see Huffman et al. in Peptides: Chemistry and Biology, G.R. Marshall ed., ESCOM Publisher: Leiden, Netherlands, 1988), substituted gama lactam rings (Garvey et al. in Peptides: Chemistry and Biology, G.R. Marshall ed., ESCOM 15. Publisher: Leiden, Netherlands, 1988), keto-methylene pseudopeptides (Ewenson et al. (1986) J Med Chem 29:295; and Ewenson et al. in Peptides: Structure and Function (Proceedings of the 9th American Peptide Symposium) Pierce Chemical Co. Rockland, IL, 1985), b-turn dipeptide cores (Nagai et al. (1985) Tetrahedron Lett 26:647; and Sato et al. (1986) J Chem Soc Perkin Trans 1:1231), and b-aminoalcohols (Gordon et al. (1985) 20 Biochem Biophys Res Commun 126:419; and et al. (1986) Biochem Biophys Res Commun 134:71).

## VACCINE FORMULATIONS FOR M. CATARRHALIS NUCLEIC ACIDS AND POLYPEPTIDES

This invention also features vaccine compositions for protection against infection by 
M. catarrhalis or for treatment of M. catarrhalis infection. In one embodiment, the vaccine 
compositions contain one or more immunogenic components such as a surface protein from 
M. catarrhalis, or portion thereof, and a pharmaceutically acceptable carrier. Nucleic acids 
within the scope of the invention are exemplified by the nucleic acids of the invention 
contained in the Sequence Listing which encode M. catarrhalis surface proteins. Any

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nucleic acid encoding an immunogenic *M. catarrhalis* protein, or portion thereof, which is capable of expression in a cell, can be used in the present invention. These vaccines have therapeutic and prophylactic utilities.

One aspect of the invention provides a vaccine composition for protection against infection by *M. catarrhalis* which contains at least one immunogenic fragment of an *M. catarrhalis* protein and a pharmaceutically acceptable carrier. Preferred fragments include peptides of at least about 10 amino acid residues in length, preferably about 10-20 amino acid residues in length, and more preferably about 12-16 amino acid residues in length.

Immunogenic components of the invention can be obtained, for example, by screening polypeptides recombinantly produced from the corresponding fragment of the nucleic acid encoding the full-length *M. catarrhalis* protein. In addition, fragments can be chemically synthesized using techniques known in the art such as conventional Merrifield solid phase f-Moc or t-Boc chemistry.

In one embodiment, immunogenic components are identified by the ability of the peptide to stimulate T cells. Peptides which stimulate T cells, as determined by, for example, T cell proliferation or cytokine secretion are defined herein as comprising at least one T cell epitope. T cell epitopes are believed to be involved in initiation and perpetuation of the immune response to the protein allergen which is responsible for the clinical symptoms of allergy. These T cell epitopes are thought to trigger early events at the level of the T helper cell by binding to an appropriate HLA molecule on the surface of an antigen presenting cell, thereby stimulating the T cell subpopulation with the relevant T cell receptor for the epitope. These events lead to T cell proliferation, lymphokine secretion, local inflammatory reactions, recruitment of additional immune cells to the site of antigen/T cell interaction, and activation of the B cell cascade, leading to the production of antibodies. A T cell epitope is the basic element, or smallest unit of recognition by a T cell receptor, where the epitope comprises amino acids essential to receptor recognition (e.g., approximately 6 or 7 amino acid residues). Amino acid sequences which mimic those of the T cell epitopes are within the scope of this invention.

Screening immunogenic components can be accomplished using one or more of several different assays. For example, *in vitro*, peptide T cell stimulatory activity is assayed

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by contacting a peptide known or suspected of being immunogenic with an antigen presenting cell which presents appropriate MHC molecules in a T cell culture. Presentation of an immunogenic *M. catarrhalis* peptide in association with appropriate MHC molecules to T cells in conjunction with the necessary co-stimulation has the effect of transmitting a signal to the T cell that induces the production of increased levels of cytokines, particularly of interleukin-2 and interleukin-4. The culture supernatant can be obtained and assayed for interleukin-2 or other known cytokines. For example, any one of several conventional assays for interleukin-2 can be employed, such as the assay described in *Proc. Natl. Acad. Sci USA*, 86: 1333 (1989) the pertinent portions of which are incorporated herein by reference. A kit for an assay for the production of interferon is also available from Genzyme Corporation (Cambridge, MA).

Alternatively, a common assay for T cell proliferation entails measuring tritiated thymidine incorporation. The proliferation of T cells can be measured *in vitro* by determining the amount of <sup>3</sup>H-labeled thymidine incorporated into the replicating DNA of cultured cells. Therefore, the rate of DNA synthesis and, in turn, the rate of cell division can be quantified.

Vaccine compositions of the invention containing immunogenic components (e.g., *M. catarrhalis* polypeptide or fragment thereof or nucleic acid encoding an *M. catarrhalis* polypeptide or fragment thereof) preferably include a pharmaceutically acceptable carrier. The term "pharmaceutically acceptable carrier" refers to a carrier that does not cause an allergic reaction or other untoward effect in patients to whom it is administered. Suitable pharmaceutically acceptable carriers include, for example, one or more of water, saline, phosphate buffered saline, dextrose, glycerol, ethanol and the like, as well as combinations thereof. Pharmaceutically acceptable carriers may further comprise minor amounts of auxiliary substances such as wetting or emulsifying agents, preservatives or buffers, which enhance the shelf life or effectiveness of the antibody. For vaccines of the invention containing *M. catarrhalis* polypeptides, the polypeptide is co-administered with a suitable adjuvant.

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It will be apparent to those of skill in the art that the therapeutically effective amount of DNA or protein of this invention will depend, *inter alia*, upon the administration schedule, the unit dose of antibody administered, whether the protein or DNA is administered in combination with other therapeutic agents, the immune status and health of the patient, and the therapeutic activity of the particular protein or DNA.

Vaccine compositions are conventionally administered parenterally, e.g., by injection, either subcutaneously or intramuscularly. Methods for intramuscular immunization are described by Wolff et al. (1990) Science 247: 1465-1468 and by Sedegah et al. (1994) Immunology 91: 9866-9870. Other modes of administration include oral and pulmonary formulations, suppositories, and transdermal applications. Oral immunization is preferred over parenteral methods for inducing protection against infection by M. catarrhalis. Cain et. al. (1993) Vaccine 11: 637-642. Oral formulations include such normally employed excipients as, for example, pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate, and the like.

The vaccine compositions of the invention can include an adjuvant, including, but not limited to aluminum hydroxide; N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP); N-acetyl-nor-muramyl-L-alanyl-D-isoglutamine (CGP 11637, referred to as nor-MDP); N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphos-phoryloxy)-ethylamine (CGP 19835A, referred to a MTP-PE); RIBI, which contains three components from bacteria; monophosphoryl lipid A; trehalose dimycoloate; cell wall skeleton (MPL + TDM + CWS) in a 2% squalene/Tween 80 emulsion; and cholera toxin. Others which may be used are non-toxic derivatives of cholera toxin, including its B subunit, and/or conjugates or genetically engineered fusions of the *M. catarrhalis* polypeptide with cholera toxin or its B subunit, procholeragenoid, fungal polysaccharides, including schizophyllan, muramyl dipeptide, muramyl dipeptide derivatives, phorbol esters, labile toxin of *E. coli*, non-*M. catarrhalis* bacterial lysates, block polymers or saponins.

Other suitable delivery methods include biodegradable microcapsules or immunostimulating complexes (ISCOMs), cochleates, or liposomes, genetically engineered attenuated live vectors such as viruses or bacteria, and recombinant (chimeric) virus-like

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particles, e.g., bluetongue. The amount of adjuvant employed will depend on the type of adjuvant used. For example, when the mucosal adjuvant is cholera toxin, it is suitably used in an amount of 5 mg to 50 mg, for example 10 mg to 35 mg. When used in the form of microcapsules, the amount used will depend on the amount employed in the matrix of the microcapsule to achieve the desired dosage. The determination of this amount is within the skill of a person of ordinary skill in the art.

Carrier systems in humans may include enteric release capsules protecting the antigen from the acidic environment of the stomach, and including *M. catarrhalis* polypeptide in an insoluble form as fusion proteins. Suitable carriers for the vaccines of the invention are enteric coated capsules and polylactide-glycolide microspheres. Suitable diluents are 0.2 N NaHCO<sub>3</sub> and/or saline.

Vaccines of the invention can be administered as a primary prophylactic agent in adults or in children, as a secondary prevention, after successful eradication of M. catarrhalis in an infected host, or as a therapeutic agent in the aim to induce an immune response in a susceptible host to prevent infection by M. catarrhalis. The vaccines of the invention are administered in amounts readily determined by persons of ordinary skill in the art. Thus, for adults a suitable dosage will be in the range of 10 mg to 10 g, preferably 10 mg to 100 mg. A suitable dosage for adults will also be in the range of 5 mg to 500 mg. Similar dosage ranges will be applicable for children. Those skilled in the art will recognize that the optimal dose may be more or less depending upon the patient's body weight, disease, the route of administration, and other factors. Those skilled in the art will also recognize that appropriate dosage levels can be obtained based on results with known oral vaccines such as, for example, a vaccine based on an E. coli lysate (6 mg dose daily up to total of 540 mg) and with an enterotoxigenic E. coli purified antigen (4 doses of 1 mg) (Schulman et al., J. Urol. 150:917-921 (1993); Boedecker et al., American Gastroenterological Assoc. 999:A-222 (1993)). The number of doses will depend upon the disease, the formulation, and efficacy data from clinical trials. Without intending any limitation as to the course of treatment, the treatment can be administered over 3 to 8 doses for a primary immunization schedule over 1 month (Boedeker, American Gastroenterological Assoc. 888:: A-222 (1993)).

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In a preferred embodiment, a vaccine composition of the invention can be based on a killed whole *E. coli* preparation with an immunogenic fragment of an *M. catarrhalis* protein of the invention expressed on its surface or it can be based on an *E. coli* lysate, wherein the killed *E. coli* acts as a carrier or an adjuvant.

It will be apparent to those skilled in the art that some of the vaccine compositions of the invention are useful only for preventing M. catarrhalis infection, some are useful only for treating M. catarrhalis infection, and some are useful for both preventing and treating M. catarrhalis infection. In a preferred embodiment, the vaccine composition of the invention provides protection against M. catarrhalis infection by stimulating humoral and/or cell-mediated immunity against M. catarrhalis. It should be understood that amelioration of any of the symptoms of M. catarrhalis infection is a desirable clinical goal, including a lessening of the dosage of medication used to treat M. catarrhalis -caused disease, or an increase in the production of antibodies in the serum or mucous of patients.

### 15 ANTIBODIES REACTIVE WITH M. CATARRHALIS POLYPEPTIDES

The invention also includes antibodies specifically reactive with the subject *M. catarrhalis* polypeptide. Anti-protein/anti-peptide antisera or monoclonal antibodies can be made by standard protocols (See, for example, *Antibodies:* A *Laboratory Manual* ed. by Harlow and Lane (Cold Spring Harbor Press: 1988)). A mammal such as a mouse, a hamster or rabbit can be immunized with an immunogenic form of the peptide. Techniques for conferring immunogenicity on a protein or peptide include conjugation to carriers or other techniques well known in the art. An immunogenic portion of the subject *M. catarrhalis* polypeptide can be administered in the presence of adjuvant. The progress of immunization can be monitored by detection of antibody titers in plasma or serum. Standard ELISA or other immunoassays can be used with the immunogen as antigen to assess the levels of antibodies.

In a preferred embodiment, the subject antibodies are immunospecific for antigenic determinants of the *M. catarrhalis* polypeptides of the invention, e.g. antigenic determinants of a polypeptide of the invention contained in the Sequence Listing, or a closely related human or non-human mammalian homolog (e.g., 90% homologous, more preferably at least

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about 95% homologous). In yet a further preferred embodiment of the invention, the anti-M. catarrhalis antibodies do not substantially cross react (i.e., react specifically) with a protein which is for example, less than 80% percent homologous to a sequence of the invention contained in the Sequence Listing. By "not substantially cross react", it is meant that the antibody has a binding affinity for a non-homologous protein which is less than 10 percent, more preferably less than 5 percent, and even more preferably less than 1 percent, of the binding affinity for a protein of the invention contained in the Sequence Listing. In a most preferred embodiment, there is no cross-reactivity between bacterial and mammalian antigens.

10 The term antibody as used herein is intended to include fragments thereof which are also specifically reactive with M. catarrhalis polypeptides. Antibodies can be fragmented using conventional techniques and the fragments screened for utility in the same manner as described above for whole antibodies. For example, F(ab')2 fragments can be generated by treating antibody with pepsin. The resulting  $F(ab')_2$  fragment can be treated to reduce disulfide bridges to produce Fab' fragments. The antibody of the invention is further intended to include bispecific and chimeric molecules having an anti-M. catarrhalis portion.

Both monoclonal and polyclonal antibodies (Ab) directed against M. catarrhalis polypeptides or M. catarrhalis polypeptide variants, and antibody fragments such as Fab' and F(ab')2, can be used to block the action of M. catarrhalis polypeptide and allow the study of the role of a particular M. catarrhalis polypeptide of the invention in aberrant or unwanted intracellular signaling, as well as the normal cellular function of the M. catarrhalis and by microinjection of anti-M. catarrhalis polypeptide antibodies of the present invention.

Antibodies which specifically bind M. catarrhalis epitopes can also be used in immunohistochemical staining of tissue samples in order to evaluate the abundance and pattern of expression of M. catarrhalis antigens. Anti-M. catarrhalis polypeptide antibodies can be used diagnostically in immuno-precipitation and immuno-blotting to detect and evaluate M. catarrhalis levels in tissue or bodily fluid as part of a clinical testing procedure. Likewise, the ability to monitor M. catarrhalis polypeptide levels in an individual can allow

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determination of the efficacy of a given treatment regimen for an individual afflicted with such a disorder. The level of an *M. catarrhalis* polypeptide can be measured in cells found in bodily fluid, such as in urine samples or can be measured in tissue, such as produced by gastric biopsy. Diagnostic assays using anti-*M. catarrhalis* antibodies can include, for example, immunoassays designed to aid in early diagnosis of *M. catarrhalis* infections. The present invention can also be used as a method of detecting antibodies contained in samples from individuals infected by this bacterium using specific *M. catarrhalis* antigens.

Another application of anti-*M. catarrhalis* polypeptide antibodies of the invention is in the immunological screening of cDNA libraries constructed in expression vectors such as λgt11, λgt18-23, λZAP, and λORF8. Messenger libraries of this type, having coding sequences inserted in the correct reading frame and orientation, can produce fusion proteins. For instance, λgt11 will produce fusion proteins whose amino termini consist of β-galactosidase amino acid sequences and whose carboxy termini consist of a foreign polypeptide. Antigenic epitopes of a subject *M. catarrhalis* polypeptide can then be detected with antibodies, as, for example, reacting nitrocellulose filters lifted from infected plates with anti-*M. catarrhalis* polypeptide antibodies. Phage, scored by this assay, can then be isolated from the infected plate. Thus, the presence of *M. catarrhalis* gene homologs can be detected and cloned from other species, and alternate isoforms (including splicing variants) can be detected and cloned.

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## KITS CONTAINING NUCLEIC ACIDS, POLYPEPTIDES OR ANTIBODIES OF THE INVENTION

The nucleic acid, polypeptides and antibodies of the invention can be combined with other reagents and articles to form kits. Kits for diagnostic purposes typically comprise the nucleic acid, polypeptides or antibodies in vials or other suitable vessels. Kits typically comprise other reagents for performing hybridization reactions, polymerase chain reactions (PCR), or for reconstitution of lyophilized components, such as aqueous media, salts, buffers, and the like. Kits may also comprise reagents for sample processing such as detergents, chaotropic salts and the like. Kits may also comprise immobilization means such as particles, supports, wells, dipsticks and the like. Kits may also comprise labeling means

such as dyes, developing reagents, radioisotopes, fluorescent agents, luminescent or chemiluminescent agents, enzymes, intercalating agents and the like. With the nucleic acid and amino acid sequence information provided herein, individuals skilled in art can readily assemble kits to serve their particular purpose. Kits further can include instructions for use.

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#### BIO CHIP TECHNOLOGY

The nucleic acid sequence of the present invention may be used to detect *M.* catarrhalis or other species of *Moraxella* acid sequence using bio chip technology. Bio chips containing arrays of nucleic acid sequence can also be used to measure expression of genes of *M. catarrhalis* or other species of *Moraxella*. For example, to diagnose a patient with a *M. catarrhalis* or other *Moraxella* infection, a sample from a human or animal can be used as a probe on a bio chip containing an array of nucleic acid sequence from the present invention. In addition, a sample from a disease state can be compared to a sample from a non-disease state which would help identify a gene that is up-regulated or expressed in the disease state. This would provide valuable insight as to the mechanism by which the disease manifests. Changes in gene expression can also be used to identify critical pathways involved in drug transport or metabolism, and may enable the identification of novel targets involved in virulence or host cell interactions involved in maintenance of an infection. Procedures using such techniques have been described by Brown *et al.*, 1995, *Science* 270: 467-470.

Bio chips can also be used to monitor the genetic changes of potential therapeutic compounds including, deletions, insertions or mismatches. Once the therapeutic is added to the patient, changes to the genetic sequence can be evaluated for its efficacy. In addition, the nucleic acid sequence of the present invention can be used to determine essential genes in cell cycling. As described in Iyer *et al.*, 1999 (*Science*, 283:83-87) genes essential in the cell cycle can be identified using bio chips. Furthermore, the present invention provides nucleic acid sequence which can be used with bio chip technology to understand regulatory networks in bacteria, measure the response to environmental signals or drugs as in drug screening, and study virulence induction. (Mons *et al.*, 1998, *Nature Biotechnology*, 16: 45-48. Patents teaching this technology include U.S. Patents 5445934, 5744305, and 5800992.

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### DRUG SCREENING ASSAYS USING M. CATARRHALIS POLYPEPTIDES

By making available purified and recombinant *M. catarrhalis* polypeptides, the present invention provides assays which can be used to screen for drugs which are either agonists or antagonists of the normal cellular function, in this case, of the subject *M. catarrhalis* polypeptides, or of their role in intracellular signaling. Such inhibitors or potentiators may be useful as new therapeutic agents to combat *M. catarrhalis* infections in humans. A variety of assay formats will suffice and, in light of the present inventions, will be comprehended by the person skilled in the art.

In many drug screening programs which test libraries of compounds and natural extracts, high throughput assays are desirable in order to maximize the number of compounds surveyed in a given period of time. Assays which are performed in cell-free systems, such as may be derived with purified or semi-purified proteins, are often preferred as "primary" screens in that they can be generated to permit rapid development and relatively easy detection of an alteration in a molecular target which is mediated by a test compound. Moreover, the effects of cellular toxicity and/or bioavailability of the test compound can be generally ignored in the *in vitro* system, the assay instead being focused primarily on the effect of the drug on the molecular target as may be manifest in an alteration of binding affinity with other proteins or change in enzymatic properties of the molecular target. Accordingly, in an exemplary screening assay of the present invention, the compound of interest is contacted with an isolated and purified *M. catarrhalis* polypeptide.

Screening assays can be constructed *in vitro* with a purified *M. catarrhalis* polypeptide or fragment thereof, such as an *M. catarrhalis* polypeptide having enzymatic activity, such that the activity of the polypeptide produces a detectable reaction product. The efficacy of the compound can be assessed by generating dose response curves from data obtained using various concentrations of the test compound. Moreover, a control assay can also be performed to provide a baseline for comparison. Suitable products include those with distinctive absorption, fluorescence, or chemi-luminescence properties, for example, because detection may be easily automated. A variety of synthetic or naturally occurring compounds can be tested in the assay to identify those which inhibit or potentiate the activity

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of the *M. catarrhalis* polypeptide. Some of these active compounds may directly, or with chemical alterations to promote membrane permeability or solubility, also inhibit or potentiate the same activity (e.g., enzymatic activity) in whole, live *M. catarrhalis* cells.

#### 5 OVEREXPRESSION ASSAYS

Overexpression assays are based on the premise that overproduction of a protein would lead to a higher level of resistance to compounds that selectively interfere with the function of that protein. Overexpression assays may be used to identify compounds that interfere with the function of virtually any type of protein, including without limitation enzymes, receptors, DNA- or RNA-binding proteins, or any proteins that are directly or indirectly involved in regulating cell growth.

Typically, two bacterial strains are constructed. One contains a single copy of the gene of interest, and a second contains several copies of the same gene. Identification of useful inhibitory compounds of this type of assay is based on a comparison of the activity of a test compound in inhibiting growth and/or viability of the two strains. The method involves constructing a nucleic acid vector that directs high level expression of a particular target nucleic acid. The vectors are then transformed into host cells in single or multiple copies to produce strains that express low to moderate and high levels of protein encoding by the target sequence (strain A and B, respectively). Nucleic acid comprising sequences encoding the target gene can, of course, be directly integrated into the host cell.

Large numbers of compounds (or crude substances which may contain active compounds) are screened for their effect on the growth of the two strains. Agents which interfere with an unrelated target equally inhibit the growth of both strains. Agents which interfere with the function of the target at high concentration should inhibit the growth of both strains. It should be possible, however, to titrate out the inhibitory effect of the compound in the overexpressing strain. That is, if the compound is affecting the particular target that is being tested, it should be possible to inhibit the growth of strain A at a concentration of the compound that allows strain B to grow.

Alternatively, a bacterial strain is constructed that contains the gene of interest under the control of an inducible promoter. Identification of useful inhibitory agents using this

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type of assay is based on a comparison of the activity of a test compound in inhibiting growth and/or viability of this strain under both inducing and non-inducing conditions. The method involves constructing a nucleic acid vector that directs high-level expression of a particular target nucleic acid. The vector is then transformed into host cells that are grown under both non-inducing and inducing conditions (conditions A and B, respectively).

Large numbers of compounds (or crude substances which may contain active compounds) are screened for their effect on growth under these two conditions. Agents that interfere with the function of the target should inhibit growth under both conditions. It should be possible, however, to titrate out the inhibitory effect of the compound in the overexpressing strain. That is, if the compound is affecting the particular target that is being tested, it should be possible to inhibit growth under condition A at a concentration that allows the strain to grow under condition B.

### LIGAND-BINDING ASSAYS

Many of the targets according to the invention have functions that have not yet been identified. Ligand-binding assays are useful to identify inhibitor compounds that interfere with the function of a particular target, even when that function is unknown. These assays are designed to detect binding of test compounds to particular targets. The detection may involve direct measurement of binding. Alternatively, indirect indications of binding may involve stabilization of protein structure or disruption of a biological function. Non-limiting examples of useful ligand-binding assays are detailed below.

A useful method for the detection and isolation of binding proteins is the Biomolecular Interaction Assay (BIAcore) system developed by Pharmacia Biosensor and described in the manufacturer's protocol (LKB Pharmacia, Sweden). The BIAcore system uses an affinity purified anti-GST antibody to immobilize GST-fusion proteins onto a sensor chip. The sensor utilizes surface plasmon resonance which is an optical phenomenon that detects changes in refractive indices. In accordance with the practice of the invention, a protein of interest is coated onto a chip and test compounds are passed over the chip. Binding is detected by a change in the refractive index (surface plasmon resonance).

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A different type of ligand-binding assay involves scintillation proximity assays (SPA, described in U.S. Patent No. 4,568,649).

Another type of ligand binding assay, also undergoing development, is based on the fact that proteins containing mitochondrial targeting signals are imported into isolated mitochondria in vitro (Hurt et al., 1985, Embo J. 4:2061-2068; Eilers and Schatz, Nature, 1986, 322:228-231). In a mitochondrial import assay, expression vectors are constructed in which nucleic acids encoding particular target proteins are inserted downstream of sequences encoding mitochondrial import signals. The chimeric proteins are synthesized and tested for their ability to be imported into isolated mitochondria in the absence and presence of test compounds. A test compound that binds to the target protein should inhibit its uptake into isolated mitochondria in vitro.

Another ligand-binding assay is the yeast two-hybrid system (Fields and Song, 1989, Nature 340:245-246). The yeast two-hybrid system takes advantage of the properties of the GAL4 protein of the yeast Saccharomyces cerevisiae. The GAL4 protein is a transcriptional activator required for the expression of genes encoding enzymes of galactose utilization. This protein consists of two separable and functionally essential domains: an N-terminal domain which binds to specific DNA sequences (UASG); and a C-terminal domain containing acidic regions, which is necessary to activate transcription. The native GAL4 protein, containing both domains, is a potent activator of transcription when yeast are grownon galactose media. The N-terminal domain binds to DNA in a sequence-specific manner but is unable to activate transcription. The C-terminal domain contains the activating regions but cannot activate transcription because it fails to be localized to UASG. In the twohybrid system, a system of two hybrid proteins containing parts of GAL4: (1) a GAL4 DNA-binding domain fused to a protein 'X' and (2) a GAL4 activation region fused to a protein 'Y'. If X and Y can form a protein-protein complex and reconstitute proximity of the GAL4 domains, transcription of a gene regulated by UASG occurs. Creation of two hybrid proteins, each containing one of the interacting proteins X and Y, allows the activation region of UAS<sub>G</sub> to be brought to its normal site of action.

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The binding assay described in Fodor *et al.*, 1991, *Science* 251:767-773, which involves testing the binding affinity of test compounds for a plurality of defined polymers synthesized on a solid substrate, may also be useful.

Compounds which bind to the polypeptides of the invention are potentially useful as antibacterial agents for use in therapeutic compositions.

Pharmaceutical formulations suitable for antibacterial therapy comprise the antibacterial agent in conjunction with one or more biologically acceptable carriers. Suitable biologically acceptable carriers include, but are not limited to, phosphate-buffered saline, saline, deionized water, or the like. Preferred biologically acceptable carriers are physiologically or pharmaceutically acceptable carriers.

The antibacterial compositions include an antibacterial effective amount of active agent. Antibacterial effective amounts are those quantities of the antibacterial agents of the present invention that afford prophylactic protection against bacterial infections or which result in amelioration or cure of an existing bacterial infection. This antibacterial effective amount will depend upon the agent, the location and nature of the infection, and the particular host. The amount can be determined by experimentation known in the art, such as by establishing a matrix of dosages and frequencies and comparing a group of experimental units or subjects to each point in the matrix.

The antibacterial active agents or compositions can be formed into dosage unit forms, such as for example, creams, ointments, lotions, powders, liquids, tablets, capsules, suppositories, sprays, aerosols or the like. If the antibacterial composition is formulated into a dosage unit form, the dosage unit form may contain an antibacterial effective amount of active agent. Alternatively, the dosage unit form may include less than such an amount if multiple dosage unit forms or multiple dosages are to be used to administer a total dosage of the active agent. Dosage unit forms can include, in addition, one or more excipient(s), diluent(s), disintegrant(s), lubricant(s), plasticizer(s), colorant(s), dosage vehicle(s), absorption enhancer(s), stabilizer(s), bactericide(s), or the like.

For general information concerning formulations, see, e.g., Gilman et al. (eds.), 1990, Goodman and Gilman's: The Pharmacological Basis of Therapeutics, 8th ed., Pergamon Press; and Remington's Pharmaceutical Sciences, 17th ed., 1990, Mack Publishing Co.,

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Easton, PA; Avis et al. (eds.), 1993, Pharmaceutical Dosage Forms: Parenteral Medications, Dekker, New York; Lieberman et al (eds.), 1990, Pharmaceutical Dosage Forms: Disperse Systems, Dekker, New York.

The antibacterial agents and compositions of the present invention are useful for preventing or treating *M. catarrhalis* infections. Infection prevention methods incorporate a prophylactically effective amount of an antibacterial agent or composition. A prophylactically effective amount is an amount effective to prevent *M. catarrhalis* infection and will depend upon the specific bacterial strain, the agent, and the host. These amounts can be determined experimentally by methods known in the art and as described above.

M. catarrhalis infection treatment methods incorporate a therapeutically effective amount of an antibacterial agent or composition. A therapeutically effective amount is an amount sufficient to ameliorate or eliminate the infection. The prophylactically and/or therapeutically effective amounts can be administered in one administration or over repeated administrations. Therapeutic administration can be followed by prophylactic administration, once the initial bacterial infection has been resolved.

The antibacterial agents and compositions can be administered topically or systemically. Topical application is typically achieved by administration of creams, ointments, lotions, or sprays as described above. Systemic administration includes both oral and parental routes. Parental routes include, without limitation, subcutaneous, intramuscular, intraperitoneal, intravenous, transdermal, inhalation and intranasal administration.

#### EXEMPLIFICATION

### 25 CLONING AND SEQUENCING M. CATARRHALIS GENOMIC SEQUENCE

This invention provides nucleotide sequences of the genome of *M. catarrhalis* which thus comprises a DNA sequence library of *M. catarrhalis* genomic DNA. The invention also provides nucleotide sequences of two naturally occurring plasmids in *M. catarrhalis*. The detailed description that follows provides nucleotide sequences of *M. catarrhalis*, and also describes how the sequences were obtained and how ORFs (Open Reading Frames) and

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protein-coding sequences can be identified. Also described are methods of using the disclosed *M. catarrhalis* sequences in methods including diagnostic and therapeutic applications. Furthermore, the library can be used as a database for identification and comparison of medically important sequences in this and other strains of *M. catarrhalis* as well as other species of *Moraxella*.

Chromosomal DNA from strain 98-4362. of *M. catarrhalis*, was isolated using a protocol described by Storrs, et al.(*J. Bacteriol*. 173: 4347-4352 (1991). The only exception to this protocol was that lysostaphin (120 U/ml) was used instead of lysozyme. The genomic DNA prep involved a lysozyme:lysostaphin digestion, sodium dodecyl sulfate lysis, Proteinase K and RNase treatment, phenol:chloroform extraction, and sodium acetate precipitation, followed by the CsCl gradient to remove the plasmid.

In the construction of both libraries, genomic *M. catarrhalis* DNA was hydrodynamically sheared in an HPLC and then separated on a standard 1% agarose gel. A fraction corresponding to 2000-3000 bp in length was excised from the gel and purified by the GeneClean procedure (Bio101, Inc.).

The purified DNA fragments were then blunt-ended using T4 DNA polymerase. The healed DNA was then ligated to unique BstXI-linker adapters (5'-GTCTTCACCACGGGG-3' and 5'-GTGGTGAAGAC-3' in 100-1000 fold molar excess). These linkers are complimentary to the BstXI-cut pGTC vector, while the overhang is not self-complimentary. Therefore, the linkers will not concatermerize nor will the cut-vector religate itself easily. The linker-adapted inserts were separated from the unincorporated linkers on a 1% agarose gel and purified using GeneClean. The linker-adapted inserts were then ligated to *Bst*XI-cut vector to construct a "shotgun" sublclone libraries.

Only major modifications to the protocols are highlighted. Briefly, the library was then transformed into DH5á competent cells (Gibco/BRL, DH5a transformation protocol). It was assessed by plating onto antibiotic plates containing ampicillin and IPTG/Xgal. The plates were incubated overnight at 37°C. Transformants were then used for plating of clones and picking for sequencing. The cultures were grown overnight at 37°C. DNA was purified using a silica bead DNA preparation (Engelstein, 1996) method. In this manner, 25 µg of DNA was obtained per clone.

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These purified DNA samples were then sequenced using primarily ABI dyeterminator chemistry. All subsequent steps were based on sequencing by ABI377 automated DNA sequencing methods. The ABI dye terminator sequence reads were run on ABI377 machines and the data was transferred to UNIX machines following lane tracking of the gels. Base calls and quality scores were determined using the program PHRED (Ewing et al., 1998, Genome Res. 8: 175-185; Ewing and Green, 1998, Genome Res. 8: 685-734). Reads were assembled using PHRAP (P. Green, Abstracts of DOE Human Genome Program Contractor-Grantee Workshop V, Jan. 1996, p.157) with default program parameters and quality scores.

Finishing followed the initial assembly. Missing mates (sequences from clones that only gave reads from one end of the *Moraxella* DNA inserted in the plasmid) were identified and sequenced with ABI technology to allow the identification of additional overlapping contigs.

End-sequencing of randomly picked genomic lambda was also performed. 15 . Sequencing of both sides was done for all lambda sequences. The lambda library backbone helped to verify the integrity of the assembly and allowed closure of some of the physical gaps. Primers for walking off the ends of contigs would be selected using pick primer (a GTC program) near the ends of the clones to facilitate gap closure. These walks can be sequenced using the selected clones and primers. These data are then reassembled with PHRAP. Additional sequencing using PCR-generated templates and screened and/or unscreened lambda templates can be done in addition.

Additional templates for the physical gaps were obtained through PCR using primers designed from the ends of the contigs. These templates were then used in sequencing reactions to close the gaps.

25 Contigs were ordered by aligning identified M. catarrhalis genes to the published physical maps. Order was confirmed by PCR. The final chromosomal assembly included 119 contigs.

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To identify *M. catarrhalis* polypeptides the complete genomic sequence of *M. catarrhalis* were analyzed essentially as follows: First, all possible stop-to- stop open reading frames (ORFs) greater than 180 nucleotides in all six reading frames were translated into amino acid sequences. Second, the identified ORFs were analyzed for homology to known (archeabacter, prokaryotic and eukaryotic) protein sequences. Third, the coding potential of non-homologous sequences were evaluated with the program GENEMARKTM (Borodovsky and McIninch, 1993, Comp. Chem. 17:123)

## IDENTIFICATION, CLONING AND EXPRESSION OF M. CATARRHALIS NUCLEIC ACIDS

Expression and purification of the *M. catarrhalis* polypeptides of the invention can be performed essentially as outlined below.

To facilitate the cloning, expression and purification of membrane and secreted proteins from *M. catarrhalis*, a gene expression system, such as the pET System (Novagen), for cloning and expression of recombinant proteins in *E. coli*, is selected. Also, a DNA sequence encoding a peptide tag, the His-Tag, is fused to the 3' end of DNA sequences of interest in order to facilitate purification of the recombinant protein products. The 3' end is selected for fusion in order to avoid alteration of any 5' terminal signal sequence.

# 20 PCR AMPLIFICATION AND CLONING OF NUCLEIC ACIDS CONTAINING ORF'S ENCODING ENZYMES

Nucleic acids chosen (for example, from the nucleic acids set forth in SEQ ID NO: 1 - SEQ ID NO: 2501 for cloning from the 98-4362. strain of *M. catarrhalis* and plasmids are prepared for amplification cloning by polymerase chain reaction (PCR). Synthetic oligonucleotide primers specific for the 5' and 3' ends of open reading frames (ORFs) are designed and purchased from GibcoBRL Life Technologies (Gaithersburg, MD, USA). All forward primers (specific for the 5' end of the sequence) are designed to include an NcoI cloning site at the extreme 5' terminus. These primers are designed to permit initiation of protein translation at a methionine residue followed by a valine residue and the coding

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sequence for the remainder of the native *M. catarrhalis* DNA sequence. All reverse primers (specific for the 3' end of any *M. catarrhalis* ORF) include a EcoRI site at the extreme 5' terminus to permit cloning of each *M. catarrhalis* sequence into the reading frame of the pET-28b. The pET-28b vector provides sequence encoding an additional 20 carboxy-terminal amino acids including six histidine residues (at the extreme C-terminus), which comprise the His-Tag.

Genomic DNA or plasmid DNA prepared from the 98-4362. strain of *M. catarrhalis* is used as the source of template DNA for PCR amplification reactions (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994). To amplify a DNA sequence containing an *M. catarrhalis* ORF, genomic DNA (50 nanograms) is introduced into a reaction vial containing 2 mM MgCl<sub>2</sub>, 1 micromolar synthetic oligonucleotide primers (forward and reverse primers) complementary to and flanking a defined *M. catarrhalis* ORF, 0.2 mM of each deoxynucleotide triphosphate; dATP, dGTP, dCTP, dTTP and 2.5 units of heat stable DNA polymerase (Amplitaq, Roche Molecular Systems, Inc., Branchburg, NJ, USA) in a final volume of 100 microliters.

Upon completion of thermal cycling reactions, each sample of amplified DNA is washed and purified using the Qiaquick Spin PCR purification kit (Qiagen, Gaithersburg, MD, USA). All amplified DNA samples are subjected to digestion with the restriction endonucleases, e.g., NcoI and EcoRI (New England BioLabs, Beverly, MA, USA)(Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994). DNA samples are then subjected to electrophoresis on 1.0 % NuSeive (FMC BioProducts, Rockland, ME USA) agarose gels. DNA is visualized by exposure to ethidium bromide and long wave uv irradiation. DNA contained in slices isolated from the agarose gel is purified using the Bio 101 GeneClean Kit protocol (Bio 101 Vista, CA, USA).

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### CLONING OF M. CATARRHALIS NUCLEIC ACIDS INTO AN EXPRESSION VECTOR

The pET-28b vector is prepared for cloning by digestion with restriction endonucleases, e.g., NcoI and EcoRI (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994). The pET-28a vector, which encodes a His-Tag

that can be fused to the 5 end of an inserted gene, is prepared by digestion with appropriate restriction endonucleases.

Following digestion, DNA inserts are cloned (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994) into the previously digested pET-28b expression vector. Products of the ligation reaction are then used to transform the BL21 strain of *E. coli* (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994) as described below.

## TRANSFORMATION OF COMPETENT BACTERIA WITH RECOMBINANT PLASMIDS

Competent bacteria, *E coli* strain BL21 or *E. coli* strain BL21(DE3), are transformed with recombinant pET expression plasmids carrying the cloned *M. catarrhalis* sequences according to standard methods (Current Protocols in Molecular, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994). Briefly, 1 microliter of ligation reaction is mixed with 50 microliters of electrocompetent cells and subjected to a high voltage pulse, after which, samples are incubated in 0.45 milliliters SOC medium (0.5% yeast extract, 2.0 % tryptone, 10 mM NaCl, 2.5 mM KCl, 10 mM MgCl2, 10 mM MgSO4 and 20, mM glucose) at 37 ©C with shaking for 1 hour. Samples are then spread on LB agar plates containing 25 microgram/ml kanamycin sulfate for growth overnight. Transformed colonies of BL21 are then picked and analyzed to evaluate cloned inserts as described below.

## IDENTIFICATION OF RECOMBINANT EXPRESSION VECTORS WITH M. CATARRHALIS NUCLEIC ACIDS

Individual BL21 clones transformed with recombinant pET-28b *M. catarrhalis* ORFs are analyzed by PCR amplification of the cloned inserts using the same forward and reverse primers, specific for each *M. catarrhalis* sequence, that were used in the original PCR amplification cloning reactions. Successful amplification verifies the integration of the *M. catarrhalis* sequences in the expression vector (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994).

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#### ISOLATION AND PREPARATION OF NUCLEIC ACIDS FROM TRANSFORMANTS

Individual clones of recombinant pET-28b vectors carrying properly cloned *M.* catarrhalis ORFs are picked and incubated in 5 mls of LB broth plus 25 microgram/ml kanamycin sulfate overnight. The following day plasmid DNA is isolated and purified using the Qiagen plasmid purification protocol (Qiagen Inc., Chatsworth, CA, USA).

#### EXPRESSION OF RECOMBINANT M. CATARRHALIS SEQUENCES IN E. COLI

The pET vector can be propagated in any *E. coli* K-12 strain e.g. HMS174, HB101, JM109, DH5, etc. for the purpose of cloning or plasmid preparation. Hosts for expression include *E. coli* strains containing a chromosomal copy of the gene for T7 RNA polymerase. These hosts are lysogens of bacteriophage DE3, a lambda derivative that carries the lacI gene, the lacUV5 promoter and the gene for T7 RNA polymerase. T7 RNA polymerase is induced by addition of isopropyl-B-D-thiogalactoside (IPTG), and the T7 RNA polymerase transcribes any target plasmid, such as pET-28b, carrying its gene of interest. Strains used include: BL21(DE3) (Studier, F.W., Rosenberg, A.H., Dunn, J.J., and Dubendorff, J.W. (1990) Meth. Enzymol. 185, 60-89).

To express recombinant *M. catarrhalis* sequences, 50 nanograms of plasmid DNA isolated as described above is used to transform competent BL21(DE3) bacteria as described above (provided by Novagen as part of the pET expression system kit). The lacZ gene (betagalactosidase) is expressed in the pET-System as described for the *M. catarrhalis* recombinant constructions. Transformed cells are cultured in SOC medium for 1 hour, and the culture is then plated on LB plates containing 25 micrograms/ml kanamycin sulfate. The following day, bacterial colonies are pooled and grown in LB medium containing kanamycin sulfate (25 micrograms/ml) to an optical density at 600 nM of 0.5 to 1.0 O.D. units, at which point, 1 millimolar IPTG was added to the culture for 3 hours to induce gene expression of the *M. catarrhalis* recombinant DNA constructions.

After induction of gene expression with IPTG, bacteria are pelleted by centrifugation in a Sorvall RC-3B centrifuge at 3500 x g for 15 minutes at 4 °C. Pellets are resuspended in 50 milliliters of cold 10 mM Tris-HCl, pH 8.0, 0.1 M NaCl and 0.1 mM EDTA (STE

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buffer). Cells are then centrifuged at 2000 x g for 20 min at 4 °C. Wet pellets are weighed and frozen at -80 °C until ready for protein purification.

A variety of methodologies known in the art can be utilized to purify the isolated proteins. (Current Protocols in Protein Science, John Wiley and Sons, Inc., J. E. Coligan et al., eds., 1995). For example, the frozen cells may be thawed, resupended in buffer and ruptured by several passages through a small volume microfluidizer (Model M-110S, Microfluidics International Corporation, Newton, MA). The resultant homogenate may be centrifuged to yield a clear supernatant (crude extract) and following filtration the crude extract may be fractionated over columns. Fractions may be monitored by absorbance at

10 OD<sub>280</sub> nm. and peak fractions may analyzed by SDS-PAGE

The concentrations of purified protein preparations may be quantified spectrophotometrically using absorbance coefficients calculated from amino acid content (Perkins, S.J. 1986 Eur. J. Biochem. 157, 169-180). Protein concentrations are also measured by the method of Bradford, M.M. (1976) Anal. Biochem. 72, 248-254, and Lowry, O.H., Rosebrough, N., Farr, A.L. & Randall, R.J. (1951) J. Biol. Chem. 193, pages 265-275, using bovine serum albumin as a standard.

SDS-polyacrylamide gels of various concentrations may be purchased from BioRad (Hercules, CA, USA), and stained with Coomassie blue. Molecular weight markers may include rabbit skeletal muscle myosin (200 kDa), *E. coli* (-galactosidase (116 kDa), rabbit muscle phosphorylase B (97.4 kDa), bovine serum albumin (66.2 kDa), ovalbumin (45 kDa), bovine carbonic anhydrase (31 kDa), soybean trypsin inhibitor (21.5 kDa), egg white lysozyme (14.4 kDa) and bovine aprotinin (6.5 kDa).

#### 25 EQUIVALENTS

Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments and methods described herein. The specific embodiments described herein are offered by way of example

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only, and the invention is to limited only by the terms of the appended claims, along with the full scope of equivalents to which such claims are entitled.

#### TABLE 2

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
2538928_c3_3	1	1921	84	255	306	3.3e-2	27
Protein name					s <u>Name</u> FA_HUMAN		Acc# P13804
Description				<del>-</del>			
ELECTRON TRANSFER	LAVOPRO	TEIN ALPH	A-SUBUNIT	PRECURS	OR (ALPHA	(-ETF)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
24035192_t2_1	2	1922	502	1509	138	1.5e-0	)5
Protein name				Locus	s Name		Acc#
icmF protein				pir:T	18341		T18341
Description							
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probab	oility
10547151_c1_2	3	1923	62	189			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT						,	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	oility
24317313_c2_2	4	1924	201	606	316	3.2e-2	:7
Protein name					Name FB_HAEIN		<u>Acc#</u> P43820
Description							
TRNA LIGASE BETA CH	AIN) (P	HERS)			· · · · · · · · · · · · · · · · · · ·		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
11911687_c2_42	5	1925	284	855	709	6.5e-7	0
Protein name 3-methyl-2-oxobutan	oate				Name 130846		<u>Acc#</u> AJ130846
Description							
Pseudomonas fluores	cens fo	IK (partia	al), panB	and pan(	(partia	l)genes	5.

ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
14334452_c2_44	6	1926	221	666	339	2.9e-35
Protein name					s Name HSDMSR	Acc# X13145
Description						<b>,</b>
Escherichia Coli pl I restriction and mo				nd hsdR	genes for	ECOR124/3 type
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16619747_c3_48	7	1927	73	222	158	1.6e-11
Protein name					s Name AV_HAEIN	<u>Acc#</u> Q57134
Description						
HYPOTHETICAL PROTEI	N HI100	8				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21657635_f3_22	8	1928	221	666	381	3.7e-35
<u>Protein name</u>					s Name BA_PSEAE	<u>Acc#</u> P95460
Description				<del> </del>		
THIOL: DISULFIDE INT	ERCHANG	E PROTEIN	DSBA PRE	CURSOR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23476431_c3_54	9	1929	181	546	123	3.8e-07
Protein name					S Name GA_HAEIN	Acc# P45076
Description						
HYPOTHETICAL PROTEI	N HI115	1				
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
23945165_c3_49	10	1930	170	513	334	3.6e-30
Protein name					Name_	Acc#
2-amino-4-hydroxy-6- dine	hydroxy	methyldihy	/dropteri	pir:C	54046	C64046
Description	" " " " " " " " " " " " " " " " " " " "			_		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24351555_c2_38	11	1931	130	393	135	1.0e-08
Protein name				Locu	s Name	Acc#
				sp:MA	ZG_HAEIN	P44723
Description						
MAZG PROTEIN HOMOLO	G			<u>,                                      </u>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2835152_t2_14	12	1932	82	249		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	NT	<u>AA</u>	Score	Probability
34178802_c2_45	13	   [1933	Length	Length 399	184	3.3e-14
		الــــــــــــــــــــــــــــــــــــ	132			
Protein name conserved hypothetic	gal gag	reted prot	ein		s Name	Acc#
HP1098	car sec.	receu prod	Jein	pir:B	6465/	B64657
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35819090_c1_31	14	1934	623	1872	689	8.5e-68
Protein name				Locus	s Name	Acc#
polynucleotide aden	ylyltra	nsferase		gp:PP	Y18131	Y18131
Description						<del></del>
Pseudomonas putida	pcnB ge	ne and par	rtial fol	K gene.		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4468761_f2_18	15	1935	61	186		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4492138_c2_43	16	1936	283	852 660	1.0e-64
Protein name				Locus Name sp:PANC_SCHPO	<u>Acc#</u> Q09673
Description					
SYNTHETASE) (PAN	TOATE ACT	IVATING EN	ZYME)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
48796812_c3_47	17	1937	84	255 138	1.8e-08
Protein name				Locus Name sp:CBF5_YEAST	Acc# P33322
Description					
5) (NUCLEOLAR PR	OTEIN CBF	5) (P64')			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5110943_c1_32	18	1938	298	897 604	8.7e-59
Protein name				Locus Name	Acc# AB033988
Description					
Shewanella viola phosphocarrier pr				nitrogen regurato eins, partialand	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5112807_c3_52	19	1939	92	279	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
26285938_f1_1	20	1940	207	624 541	4.1e-52
Protein name				Locus Name sp:YCEG HAEIN	Acc# P44720
Description					
HYPOTHETICAL PRO	TEIN HIO4	57			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
30658587_f1_2	21	1941	73	219 126	4.8e-08
Protein name				Locus Name sp:KTHY_BACSU	<u>Acc#</u> P37537
Description				<u></u>	
THYMIDYLATE KINASE	, (DTMP	KINASE)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
2767080_f1_2	22	1942	373	1122 1522	4.6e-156
Protein name				Locus Name	Acc#
				sp:EFTU_SHEPU	P33169
Description					
ELONGATION FACTOR	ru (EF-	TU)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
32110007_c2_8	23	1943	88	267 114	7.3e-07
Protein name				Locus Name	Acc#
	- BYY2 2	5'E"		pir:H71023	H71023
hypothetical protei	Ln PH14			] prr:H/1023	11/2020
Description	LN PH148			pir:H/1023	
	NTID	AAID	NT Length	AA Score	Probability
Description				AA Score	
Description ORF Name	NTID	AAID	Length	AA Score Length 183 144 Locus Name	Probability 5.5e-09 Acc#
Description  ORF Name  36329582_c1_5  Protein name	NTID	AAID	Length	AA Score Length 183	Probability 5.5e-09
Description  ORF Name  36329582_c1_5  Protein name  Description	NTID	<u>AAID</u>	Length	AA Score Length  183  Locus Name  Sp:YHA2_EIKCO	Probability 5.5e-09 Acc#
Description  ORF Name  36329582_c1_5  Protein name	NTID	<u>AAID</u>	Length 60 2 5'REGIO	AA Score Length 183 144  Locus Name sp:YHA2_EIKCO	Probability 5.5e-09 Acc#
Description  ORF Name  36329582_c1_5  Protein name  Description	NTID	<u>AAID</u>	Length	AA Score Length  183  Locus Name  Sp:YHA2_EIKCO	Probability 5.5e-09 Acc#
Description  ORF Name  36329582_c1_5  Protein name  Description  HYPOTHETICAL 66.3	NTID  24  KD PROT	AAID 1944 EIN IN HAG	Length 60 2 5'REGIO	AA Score Length 183 Locus Name Sp:YHA2_EIKCO	Probability 5.5e-09 Acc# P35649
Description  ORF Name  36329582_c1_5  Protein name  Description  HYPOTHETICAL 66.3	NTID  24  KD PROT	AAID  1944  EIN IN HAG  AAID	Length 60  2 5'REGIO NT Length	AA Score Length  183  Locus Name Sp:YHA2_EIKCO  N  AA Score Length Score	Probability  5.5e-09  Acc# P35649  Probability
Description  ORF Name  36329582_c1_5  Protein name  Description  HYPOTHETICAL 66.3 1  ORF Name  971016_f1_1	NTID  24  KD PROT	AAID  1944  EIN IN HAG  AAID	Length 60  2 5'REGIO NT Length	AA Score Length  183  Locus Name Sp:YHA2_EIKCO  AA Score Length  597  643  Locus Name	Probability  5.5e-09  Acc# P35649  Probability  6.4e-63  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
2225312_f3_18	26	1946	427	1284 416	1.9e-64
Protein name  glycerophosphoryl	diester	phosphod	iesterase	Locus Name	Acc# D75630
Description					
ORF Name	NTID	AAID	NT Length	AA Score	Probability
23457692_f1_1	27	1947	392	1179 360	1.9e-42
Protein name				Locus Name sp:RECF_PSEPO	Acc# P13456
Description					
RECF PROTEIN					
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
26042927_f3_19	28	1948	84	255	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
26750837_t1_4	29	1949	111	336 202	4.9e-16
Protein name				Locus Name	Acc#
hypothetical prote	in			pir:S76551	S76551
Description		•			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
36144675_f1_2	30	1950	525	1578 1851	6.3e-191
Protein name				Locus Name sp:GUAA_HAEIN	Acc# P44335
Description					
AMIDOTRANSFERASE)	(GMP SY	THETASE)			

ORF Name	NTID AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
4298443_f2_8	31 1951	822	2469	2597	5.6e-270
Protein name			Locus	Name	Acc#
Description			sp:GYI	RB_ECOLI	P06982:008 438
DNA GYRASE SUBUNIT	В,		,	,	
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12617627_c1_1	32 1952	128	387	650	1.2e-63
Protein name			Locus	Name	\ <u>Acc#</u>
transposase			pir:I	57760	167760
<u>Description</u>					
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34175180_c2_2	33 1953	90	273	137	1.7e-08
Protein name			Locus	Name	Acc#
transposase			gp:AB0	26428	AB026428
Description					
Methylomonas aminoi IS10-R rmpI, rmpB),		monophospl	nate path	way gene	es(rmpD, rmpA,
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16690875_f1_2	34 1954	82	249	90	0.00026
Protein name			Locus	Name	Acc#
TolR protein			gp:PPI	PAL1	X74218
Description					
Pseudomonas putida	ruvB, tolQ, tolR,	, toIA, to	olB and c	prL gene	es.
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1953953_c2_15	35 1955	534	1605	1387	9.3e-142
Protein name				Name	Acc#
Description			sp:AN]	A_NEIGO	Q02219
MAJOR OUTER MEMBRAN	NE PROTEIN PAN 1 I	PRECURSOR			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22667557_f2_6	36	1956	177	534	260	4.5e-31
Protein name					s Name DE_BACSU	<u>Acc#</u> 007573
Description						· .
HYPOTHETICAL 16.6	KD PROTE	EIN IN GL	PD-SPOVR I	NTERGENI	C REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30644217_f2_8	37	1957	83	252		
Protein name				Locu	s Name	Acc#
<u>Description</u>						
NO-HIT				•		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4881533_f2_7	38	1958	62	189		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT					,,,,	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6651712_t2_2	39	1959	271	816	611	1.6e-59
<u>Protein name</u>				Locus	s Name	Acc#
isocitrate lyase				gp:AB	004651	AB004651
Description						
Hyphomicrobium met transporter, methion	_	-		_		nicphosphate
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
14647952_f1_1	40	1960	912	2739	2108	3.7e-218
Protein name				Locus	s Name	Acc#
initiation factor	IF2-alph	ıa		gp:PV	AJ2737	AJ002737
Description						
Proteus vulgaris i	nfB gene	and part	ial nusA	and rbfA	genes.	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
15032818_c1_15	41	1961	172	519	112	4.1e-	05
Protein name hypothetical protei	n			Locus pir:G	5 Name 75410		<u>Acc#</u> G75410
Description				ţ			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score		bility
21644075_c1_14	42	1962	199	600	381	3.7e-	35
Protein name conserved hypotheti	cal pro	tein		Locus pir:F	<u> </u>		Acc# F75410
Description							
ORF Name 24650277_f1_3	NTID 43	AAID 1963	NT Length	AA Length 939	<u>Score</u>	Proba	bility 52
Protein name  Description					s Name UB_HAEIN		<u>Acc#</u> P45142
ORF Name  3332760 f2 11	NTID	AAID	NT Length	AA Length	Score	Proba	bility
Protein name	·	J		Locu	s Name		Acc#
Description							<u></u>
NO-HIT							
ORF Name 3407812 f2 9	NTID 45	<u>AAID</u>	NT Length	AA Length [507	Score	Proba	bility
Protein name				Locus	s Name FA ECOLI		Acc# P09170
Description							
RIBOSOME-BINDING FA	CTOR A	(P15B PRO	TEIN)				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4573462_c2_24	46	1966	103	312	171	2.0e-12
Protein name  conserved hypotheti	cal pro	otein		Locu:	s Name 75410	Acc# F75410
Description			•			
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
4968825_±2_5	47	1967	217	654	466	3.7e-44
Protein name					s Name SA_ECOLI	Acc# P03003
Description				1		
N UTILIZATION SUBST	ANCE PI	ROTEIN A (	NUSA PROT	EIN) (È	FACTOR)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
7070265_f1_4	48	1968	62	189		
<u>Protein name</u>				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4143942_f3_1	49	1969	319	957	164	1.1e-11
Protein name					s Name	Acc#
hypothetical protei	n b1759	) 		pir:G	64935	G64935
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
1072952_£3_19	50	1970	331	996	281	2.5e-24
Protein name					S Name G2_YEAST	Acc#
Description						P53549:Q08 718
PROBABLE 26S PROTEA	SE SUB	ONIT SUG2	(PROTEASC	MAL CAP	SUBUNIT)	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
112880_f1_4	51	1971	99	300	120	1.7e-07
Protein name hypothetical protei	n APE25	54		<u>Locu</u> pir:C	s Name 72489	<u>Acc#</u> C72489
Description						
ORF Name	NTID	AAID	NT Length	AA Length [504	Score	Probability
Protein name Description	32	] [372_]			s Name	Acc#
NO-HIT		-				
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
19532782_c2_33	53	1973	513	1542	1454	7.4e-149
Protein name					s Name PE_ACICA	Acc# P23315
Description						<del></del>
ANTHRANILATE SYNTHA	SE COME	PONENT I,				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20939567_f1_1	54	1974	138	417		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22070965_f2_11	55	1975	123	372	88	0.018
Protein name			1.5	Locus	s Name	Acc#
alaninetRNA ligas synthetase:alanyl-tR		_	ANA A	pir:D	70127	D70127
Description				_		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23839667_c1_25	56	1976	318	957	732	2.4e-72
Protein name					s Name PA_HAEIN	<u>Acc#</u> P43797
Description						
DIHYDRODIPICOLINATE	SYNTHA	ASE, (DHDP:	S)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26281300_c3_35	57	1977	119	360	257	5.1e-22
Protein name					s Name 1B_MYCTU	<u>Acc#</u> Q10514
Description						······
HYPOTHETICAL 39.6 K	D PROTE	EIN CY427.	11C			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30507291_£3_20	58	1978	174	525		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4792250_c1_26	59	1979	114	345		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5282805_c3_34	60	1980	241	726	786	4.5e-78
Protein name					s Name R7_ECOLI	Acc# P21155
Description						
(SAICAR SYNTHETASE)						

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
24020430_f2_1	61	1981	127	381	649	1.5e-63
Protein name transposase Description				Locu pir:I	s Name 67760	Acc# 167760
ORF Name 129813_f2_1 Protein name	NTID 62	<u>AAID</u>	NT Length	AA Length 381 Locu	Score s Name	Probability  Acc#
Description NO-HIT						
ORF Name 4391518_f2_4 Protein name	NTID	AAID 1983	NT Length 64		Score 108 s Name IX_HAEIN	Probability  3.2e-06  Acc# P43787
Description THIOREDOXIN-LIKE PR	ROTEIN I	HI1115				
ORF Name  4495268_f2_2  Protein name ferredoxin [3Fe-45]  Description	NTID 64	<u>AAID</u>	NT Length 110	AA Length 333 Locus pir:F	Score  512  S Name EAV	Probability  4.9e-49  Acc#  A29936:A00 218
ORF Name  4860875_f3_6  Protein name hypothetical protein Description	NTID  65 n APE24	AAID 1985	NT Length 159	AA Length 480 Locus pir:F	Score 204 S Name 72475	Probability  2.1e-16  Acc# F72475

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15677200_t2_2	66	1986	158	477	428	3.9e-40
Protein name  Description					s Name SW_ECOLI	Acc# P16702:P76 534
SULFATE TRANSPORT S	YSTEM P	ERMEASE P	-	rsw		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4490678_f1_1	67	1987	247	741	643	6.4e-63
Protein name					s Name SA ECOLI	Acc#
Description						P16676:P77 693
SULFATE TRANSPORT A	TP-BIND	ING PROTE	IN CYSA			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16054077_±3_20	68	1988	520	1563		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16495465_f1_1	69	1989	77	234	72	0.020
Protein name					s Name IE_ECOLI	Acc# P40721
Description						
HYPOTHETICAL 7.1 KD	PROTEI	N IN AROH	-NLPC INT	ERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23485750_c3_36	70	1990	68	207		
<u>Protein name</u>				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23730017_c1_24	71	1991	947	2844 278	2.2e-36
Protein name				Locus Name sp:YTFM_HAEIN	Acc# P44038
Description					
HYPOTHETICAL PROTEI	N H1069	8 PRECURS	OR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23859387_£2_14	72	1992	296	891 93	0.048
Protein name				Locus Name	Acc#
conserved hypotheti	cal pro	tein yrrB	•	pir:H69978	H69978
Description					<del></del>
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
34119002_f3_18	73	1993	444	1335 714	1.1e-69
Protein name				Locus Name	Acc#
2-acylglycerophosph acyltransferase (aas				pir:E71667	E71667
Description					
ORF Name	NTID	AAID	NT Length	AA Score	Probability
4480217_c3_35	74	1994	1675	5028 678	1.5e-79
Protein name				Locus Name sp:YTFN_HAEIN	Acc# Q57523
Description					
HYPOTHETICAL PROTEI	N H1069	96			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
12378407_c2_32	75	1995	278	834 626	4.1e-61
Protein name				Locus Name	Acc# P24223
Description				<u> </u>	J
PYRIDOXAL PHOSPHATE	BIOSYN	THETIC PR	OTEIN PDX	J	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14487952_t1_7	76	1996	72	219		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT		·				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
161402_c2_29	77	1997	61	186	59	0.018
Protein name envelope glycoprote	ein				s Name VU90070	Acc# U90070
Description						
HIV-1 strain VN16 partial cds.	trom Vie	etnam, env	elope gly	/coprotei	n V3 reg	ion(env) gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16171905_c2_28	78	1998	67	204		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22324331_f2_16	79	1999	77	234		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22463311_f3_22	80	2000	103	312		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						····

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23442503_c2_31	81	2001	346	1041	831	7.7e-83
Protein name Era	<del>-</del> ,				s Name 123492	Acc# AF123492
Description						
Pseudomonas aerugii	nosa rn	c-era-recC	operon,	complete	sequence	∍.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24412781_c3_34	82	2002	101	306		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT		<del></del>				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26570925_c2_30	83	2003	268	807	500	9.1e-48
Protein name				-	s Name	Acc#
Description				sp:RN	C_ECOLI	P05797:P06
RIBONUCLEASE III,	(RNASE	III)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
26678567_c1_24	84	2004	63	192	88	0.00042
Protein name hypothetical protein	in 29.1	<u> </u>		Locus pir:S	s Name 59084	Acc# S59084
Description				<del></del>		
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
35161562_c1_27	85	2005	212	639	103	0.0015
Protein name RecO					s Name 123492	Acc# AF123492
Description						
Pseudomonas aerugii	nosa rn	c-era-rec0	operon,	complete	sequence	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
4063308_c3_35	86	2006	607	1824	2257	5.9e-234
Protein name				Locus N		Acc# P43729
Description				<del>.</del>		
GTP-BINDING PROTEIN	LEPA					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
4100003_f3_20	87	2007	159	480	624	6.6e-61
Protein name				Locus N sp:Y882		Acc# P44068
Description						
HYPOTHETICAL PROTEI	N HIO88	32				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sc	core	Probability
7032838_c3_36	88	2008	367	1104	276	2.0e-44
Protein name signal peptidase I		•		Locus N		Acc# D64044
Description				<u> </u>		
Escherichia coli ri	bonucle	ase III a	nd other	genes, comp	olete co	ds.
ORF Name	NTID	AAID	<u>NT</u> Length	AA So	ore	Probability
9869702_£3_21	89	2009	60	183		
Protein name				Locus N	<u>lame</u>	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
10802330_f3_20	90	2010	64	195		
Protein name				Locus N	<u>fame</u>	Acc#
Description						
NO-HIT			· · · · · · · · · · · · · · · · · · ·			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12714056_c1_22	91	2011	377	1134	1472	9.1e-151
Protein name  putative formaldehy	de dehy	odrogenase		_	s Name P243941	Acc# AJ243941
Description						
Pseudomonas sp. str	ain HRI	199 partia	I vanB, f	dh, gcs,	ehyA and	ehyBgenes.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14844626_c2_34	92	2012	202	609	93	0.028
Protein name				Locus	s Name	Acc#
transcription regul	ator, T	etR family	7	pir:F	75482	F75482
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15705056_c1_24	93	2013	72	219		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
159667_c2_31	94	2014	67	204		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
30079512_f3_17	95	2015	76	231	87	0.00053
Protein name				·	s Name XS_RHIME	<u>Acc#</u> P18399
Description						
NITROGEN FIXATION P	ROTEIN	FIXS				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
35578402_£2_7	96	2016	441	1326	1116	4.8e-	113
Protein name				-	s <u>Name</u> EF_ECOLI		<u>Acc#</u> P33016
Description							
HYPOTHETICAL 49.8	KD TRANS	PORT PROT	EIN IN SE	CB-HISL	INTERGEN	C REG	ON.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
3910876_£3_16	97	2017	501	1506	742	2.1e-	73
Protein name					s Name	<del></del> -1	Acc#
Description							P77649:P76 904
HYPOTHETICAL 54.4	KD PROTE	IN IN ARO	H-NLPC IN	TERGENIC	REGION		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
5097812_c2_30	98	2018	120	363	280	1.9e-	24
Protein name				Locus	s Name		Acc#
<u>Description</u>				sp:YA	IW_ECOLI		P51025:P77 317
HYPOTHETICAL 31.4	KD PROTE	IN IN MHP	r-adhc in	TERGENIC	REGION		
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Proba	bility
5210318_t2_10	99	2019	289	870	196	1.5e-	15
Protein name		n		Locus	s Name		Acc#
hypothetical prot	ein HP086	L		pir:E	64627		E64627
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
6740877_£3_15	100	2020	406	1221	639	1.7e-	62
Protein name	: _			Locus	s Name		Acc#
stearoyl-CoA desa	iturase			gp:AF	026401		AF026401
Description							
Mucor rouxii stea	aroyl-CoA	desaturas	e (Ole1)	gene, cor	mplete co	ls.	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probabilit	У
994001_c1_23	101	2021	176	531	573	1.7e-55	
Protein name				·	s Name IG_ECOLI	Acc P330	
Description						<del></del> -	
HYPOTHETICAL 31.3	O PROTI	EIN IN FOL	E-CIRA IN	TERGENIC	REGION		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	Y
1048137_c3_65	102	2022	67	204			
Protein name				Locu	s Name	Acc	<u>#</u>
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	Y
10585925_f1_2	103	2023	73	222			
Protein name				Locus	s Name	Acc	<u>#</u>
Description					•		
NO-HIT		···					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	Υ
14885910_c2_51	104	2024	86	258	71	0.026	
Protein name				Locus	s Name	Acc	<u>#</u>
PagK				gp:AF	013775	AF01	3775
Description							
Salmonella typhimum complete cds.	rium Pag	gK (pagK),	PagM (pa	gM), and	PagO (pa	igO) genes,	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	Y
22554587_c3_57	105	2025	159	480	480	1.2e-45	
Protein name					s Name	Acc	<u>#</u>
Description				sp:SM	PB_ECOLI	P3205 011	2:P77
SMALL PROTEIN B (18	3.3 KD 1	PROTEIN)				· <del>-</del> ···	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23437838_f3_28	106	2026	725	2178	1684	3.1e-173
Protein name				·	Name	Acc# P43813
Description					-	<del></del>
)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23468813_f3_22	107	2027	309	930	294	6.2e-26
Protein name				Locus	Name	Acc#
putative permease	BitE			gp:SH	J75349	U75349
Description	,					
Serpulina hyodysen	teriae l	oit operor	, complet	e sequenc	ce.	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
234807_f1_11	108	2028	175	528	456	4.2e-43
Protein name lipopolysaccharide kdtB homolog	core b	iosynthesi	s protein		Name 72166	Acc# S72166
lipopolysaccharide	core b	iosynthesi	s protein	_		<del></del>
lipopolysaccharide kdtB homolog  Description  ORF Name	NTID	AAID	NT Length	Pir:S	Score	S72166  Probability
lipopolysaccharide kdtB homolog Description			<u>NT</u>	pir:S	72166	S72166
lipopolysaccharide kdtB homolog  Description  ORF Name	NTID	AAID	NT Length	AA Length 1083 Locus	Score	S72166  Probability
lipopolysaccharide kdtB homolog Description  ORF Name  23705040_f2_16	NTID	AAID	NT Length	AA Length [1083] Locus	Score 670	S72166  Probability  8.8e-66  Acc#
lipopolysaccharide kdtB homolog Description  ORF Name  23705040_f2_16  Protein name	NTID 109	<u>AAID</u>	NT Length 360	AA Length 1083 Locus sp:PO1	Score  670  Name  TA_HAEIN	S72166  Probability  8.8e-66  Acc#
Iipopolysaccharide kdtB homolog  Description  ORF Name  23705040_f2_16  Protein name  Description  SPERMIDINE/PUTRESC	NTID  109  . INE TRAI	AAID 2029 NSPORT ATP	NT Length 360	AA Length 1083 Locus Sp:PO1	Score  670  Name  TA_HAEIN	8.8e-66  Acc# P45171
Iipopolysaccharide kdtB homolog  Description  ORF Name  23705040_f2_16  Protein name  Description  SPERMIDINE/PUTRESC  ORF Name	NTID  109  INE TRAI	AAID 2029 NSPORT ATP	NT Length 360 P-BINDING NT Length	AA Length  Locus  Sp:POT  PROTEIN F  AA Length	Score  670  Name  TA_HAEIN  OTA  Score	Probability  8.8e-66  Acc# P45171  Probability
Iipopolysaccharide kdtB homolog  Description  ORF Name  23705040_f2_16  Protein name  Description  SPERMIDINE/PUTRESC	NTID  109  . INE TRAI	AAID 2029 NSPORT ATP	NT Length 360 -BINDING	AA Length 1083 Locus Sp:PO1	Score  670  Name  TA_HAEIN	8.8e-66  Acc# P45171
lipopolysaccharide kdtB homolog  Description  ORF Name  23705040_f2_16  Protein name  Description  SPERMIDINE/PUTRESC  ORF Name  23726687_f2_17  Protein name	NTID 109 INE TRAI	AAID 2029 NSPORT ATP AAID 2030	NT Length 360 P-BINDING NT Length 335	AA Length 1083 Locus Sp:PO1 PROTEIN F AA Length 1008 Locus	Score  670 Name CA_HAEIN  OOTA  Score  745 Name	Probability  8.8e-66  Acc# P45171  Probability  9.9e-74  Acc#
lipopolysaccharide kdtB homolog Description  ORF Name  23705040_f2_16  Protein name  Description  SPERMIDINE/PUTRESC  ORF Name  23726687_f2_17	NTID 109 INE TRAI	AAID 2029 NSPORT ATP AAID 2030	NT Length 360 P-BINDING NT Length 335	AA Length  1083  Locus  Sp:POI  PROTEIN F  AA Length  1008	Score  670 Name CA_HAEIN  OOTA  Score  745 Name	Probability  8.8e-66  Acc# P45171  Probability  9.9e-74

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23884387_c1_37	111	2031	219	660		
Protein name		•		Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
24252305_f2_19	112	2032	298	897	155	8.2e-09
Protein name				-	s Name FC_BACSU	Acc# P96680
Description						
HYPOTHETICAL 33.6	KD PROT	EIN IN CS	PC-NAP INT	ERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24797312 f1 4	113	2033	275	828	132	5.0e-06
21737312_11_1						Ł
Protein name		<b>_</b>		_	s Name	Acc#
	ein PH11	L4		Locus pir:C		Acc# C71052
Protein name	ein PH111	14		_		<del></del>
Protein name hypothetical prot	ein PH111	AAID	NT Length	_		<del></del>
Protein name hypothetical prot Description			NT	pir:C	71052	C71052
Protein name hypothetical prot Description  ORF Name	NTID	AAID	NT Length	AA Length	71052	C71052
Protein name hypothetical prot  Description  ORF Name  25901467_c3_54	NTID	AAID	NT Length	AA Length	71052 Score	C71052
Protein name hypothetical prot Description  ORF Name 25901467_c3_54 Protein name	NTID	AAID	NT Length	AA Length	71052 Score	C71052
Protein name hypothetical prot  Description  ORF Name  25901467_c3_54  Protein name  Description	NTID	AAID	NT Length	AA Length	71052 Score	C71052
Protein name hypothetical prot  Description  ORF Name  25901467_c3_54  Protein name Description  NO-HIT	NTID	AAID 2034	NT Length 88	AA Length  267  Locus	Score S Name	Probability  Acc#
Protein name hypothetical prot  Description  ORF Name  25901467_c3_54  Protein name Description  NO-HIT  ORF Name  30272051_t3_24  Protein name	NTID 114 NTID 115	AAID  AAID	NT Length 88	AA Length Locus  AA Length  Locus  Locus  Locus	Score  Score  Score  169  Name	Probability  Acc#  Probability  2.2e-11  Acc#
Protein name hypothetical prot  Description  ORF Name  25901467_c3_54  Protein name Description  NO-HIT  ORF Name  30272051_t3_24	NTID  114  NTID  115  gical	AAID    2034    AAID   2035	NT Length 88	AA Length Locus  AA Length  Locus  AA Length  747	Score  Score  Score  169  Name	Probability  Acc#  Probability  2.2e-11

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
3323802_f3_23	116	2036	286	861	232	2.3e-19
Protein name permease protein				Locus 3		<u>Acc#</u> AJ000750
Description						
Campylobacter jeji	uni malF	gene, pa	rtial.			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
35976510_f3_30	117	2037	89	270	343	4.0e-31
Protein name				Locus Dir:FEK		Acc#
Description				<b></b>		S72167:S78 121:A00210
ORF Name	NTID	AAID	NT Length	Length -	core	Probability
36383542_t1_13	118	2038	107	321	96	5.9e-05
Protein name				Locus		<u>Acc#</u>
KH type splicing i	regulator	ry proteir	1	gp:HSKH	SRP3	AF093747
Description						
Homo sapiens KH ty partial cds.	ype spli	cing regu	latory pro	tein (KHSR	P) gene	e, exon2 and
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length So	core	Probability
3923288_c1_39	119	2039	343	1032	254	1.1e-21
Protein name				Locus	Name	Acc#
probable regulator	ry prote:	in (pfoS/I	₹)	pir:E71	373	E71373
Description					•	
ORF Name	NTID	AAID	<u>NT</u> Length	Length So	core	Probability
3938393_c3_64	120	2040	218	657	726	1.0e-71
Protein name				Locus	Name	Acc#
uracil phosphoribo	syltrans	sferase, ι	ıpp	pir:A65	026	
Description						A65026:S23

ORF Name	NTID AAID	NT Length	AA Length Score	Probability
4064638_f1_3	121 2041	371	1116 152	4.7e-08
Protein name			Locus Name sp:Y131_HAEIN	Acc# P43951
Description				
HYPOTHETICAL PROTEI	N HI0131 PRECURSO	R		
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
4101568_f3_29	2042	263	792 512	4.9e-49
<u>Protein name</u>			Locus Name sp:FRP_VIBHA	<u>Acc#</u> Q56691
Description				
(NADPH-FMN OXIDORED	UCTASE)			
ORF Name	NTID AAID	NT Length	AA Length Score	Probability
682641_c1_33	123 2043	86	261 100	2.2e-05
Protein name			Locus Name	Acc#
hypothetical protei	n PH0217		pir:G71244	G71244
Description				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
10790_f3_68	124 2044	731	2196 594	9.0e-86
Ducksin nome				
Protein name			Locus Name sp:PRIM HAEIN	Acc# Q08346
Description				· · · · · · · · · · · · · · · · · · ·
				· · · · · · · · · · · · · · · · · · ·
Description	NTID AAID	<u>NT</u> Length		· · · · · · · · · · · · · · · · · · ·
Description  DNA PRIMASE,			sp:PRIM_HAEIN  AA Score	Q08346
Description  DNA PRIMASE,  ORF Name		Length 1	Sp:PRIM_HAEIN  AA Length Score	Q08346  Probability
Description  DNA PRIMASE,  ORF Name  [119012_c3_118		Length 1	Sp:PRIM_HAEIN  AA Length  1317  Locus Name	Q08346  Probability  1.1e-188  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	ore Prob	ability
12214386_c3_117	126	2046	125	378		
Protein name				Locus Na	ıme	Acc#
Description						
NO-HIT				·		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore Prob	ability
12540957_c3_121	127	2047	280	843	7.7e	-19
Protein name probable yfiH prot	ein			Locus Na	<del></del>	<u>Acc#</u> A70579
Description						
ORF Name	NTID	AAID	NT Length	AA Length	ore <u>Prob</u>	ability
12593961_f2_35	128	2048	68	207		
Protein name				Locus Na	ime	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	re Proba	ability
19532813_f3_73	129	2049	134	405	1.7e	-21
Protein name  RpsT protein				Locus Na		<u>Acc#</u> AJ002395
Description			· <del>· · · · · · · · · · · · · · · · · · </del>			l
Vibrio cholerae nhaR, hlyU, mviN, and rpsT genes.						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	re Proba	ability
209375_c1_95	130	2050	750	2250	1.3e-	-192
Protein name				Locus Na		Acc#
Description						P15716:P77 686
ATP-DEPENDENT CLP	PROTEAS	E ATP-BIN	DING SUBUN	IT CLPA		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21641077_f2_49	131	2051	199	600	132	4.3e-08
Protein name hypothetical prote	in			Locu gp:SY	s Name CSLLE	Acc# D64003:AB0
Description  Synechocystis sp.	PCC6803	complete	genome, 2	2/27, 27	55703-286	01339
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
22143827_c1_89	132	2052	250	753	246	7.5e-21
Protein name  Description					s Name V8_YEAST	Acc# P40582
	<u> </u>		1 2 1 1 1 1 1 1 1 1			
HYPOTHETICAL 26.8	KD PROT	EIN IN HYR				
ORF Name	NTID	AAID	NT Length	AA Length	Score	<u>Probability</u>
22453453_c2_104	133	2053	426	1281	492	6.4e-47
Protein name carboxyl-terminal	nvotoin	200		_	s Name	Acc#
	proceina	45E		pir:F	70369	F70369
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22831262_c1_94	134	2054	128	387	185	2.2e-14
Protein name					s Name JA_ECOLI	Acc# P75832
Description						
12.2 KD PROTEIN IN CSPD-CLPA INTERGENIC REGION						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
23632215_£2_59	135	2055	64	195		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23645875_c1_84	136	2056	605	1818	723	2.1e-71
Protein name					s <u>Name</u> DD_ECOLI	Acc# P29018:Q47
Description TRANSPORT ATP-BIND	ING PROT	EIN CYDD			· · · · ·	656:P77275
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
23875303_c2_109	137	2057	72	219		
Protein name Description				Locus	s Name	Acc#
NO-HIT		•			·	
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
24236642_c1_91	138	2058	350	1053	695	2.0e-68
Protein name					Name OD_ECOLI	Acc#
Description				<u> </u>		P33643:P77 003
(PSEUDOURIDYLATE S	YNTHASE)	(URACIL	HYDROLYAS	E)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24317757_f3_67	139	2059	368	1107	352	4.4e-32
Protein name					s Name IY PSEAE	Acc# P33641
Description						<u></u> /
HYPOTHETICAL 38.5	KD LIPOE	PROTEIN IN	PILS 5'R	EGION PRI	ECURSOR (	ORFY)
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
24318805_f2_60	140	2060	205	618	229	4.8e-19
Protein name hypothetical prote	in				Name A224767	Acc# AJ224767
<u>Description</u>				<u>ـــــــ</u>		
Acinetobacter sp.	ADP1 lor	gene and	ORFs.			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24417012_f2_52	141	2061	232	699 101	0.011
Protein name LpsB				Locus Name gp:AF193023	Acc# AF193023
Description					
Sinorhizobium meli LpsC (lpsC), and Lr		_			, LpsD(lpsD),
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
24650062_c3_119	142	2062	234	705 148	3.1e-09
Protein name hypothetical prote	in C33F	10.3		Locus Name	Acc# T15745
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
2468950_c3_123	143	2063	67	204 123	1.4e-06
Protein name				Locus Name sp:COPA HELFE	<u>Acc#</u> 032619
Description				-	
COPPER-TRANSPORTIN	G ATPAS	Ε,			
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
25391941_c2_116	144	2064	298	897	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
261635_£1_16	145	2065	217	654 603	1.1e-58
Protein name				Locus Name	Acc#
response regulator	GacA			gp:AF115381	AF115381
Description					
Pseudomonas aureofcds.	aciens	30-84 resp	onse regu	ilator GacA (gacA	) gene, complete

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31431512_f1_22	146	2066	182	549	295	4.8e-26
Protein name  bacterioferritin co	omigrato	ory protein	າ	Locus pir:F	Name 71971	Acc# F71971
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
31832188_c2_114	147	2067	440	1323	1025	2.1e-103
Protein name					s Name 90_HAEIN	<u>Acc#</u> P77868
Description						
PROBABLE CATION-TRA	MSPORTI	NG ATPASE	HI0290,			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33845302_c2_115	148	2068	288	867	653	5.6e-64
Protein name					s Name 90_HAEIN	Acc# P77868
Description						
PROBABLE CATION-TRA	NSPORT	ING ATPASE	HI0290,			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
35974750_f2_38	149	2069	261	786	603	1.1e-58
Protein name					S <u>Name</u> GI_HAEIN	Acc#
Description						Q57354:005 008
HYPOTHETICAL PROTEI	N HIOI	)5				
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4806512_c2_96	150	2070	463	1392	1501	7.7e-154
Protein name					s Name	Acc#
hypothetical protei	.n 7			pir:T	00129	T00129
Description						

ORF Name	NTID AAID	<u>NT</u> Length	AA Score	Probability
5109843_c2_99	151 2071	579	1740 291	4.3e-45
Protein name			Locus Name sp:CYDC_ECOL1	Acc# P23886
Description				
TRANSPORT ATP-BIN	DING PROTEIN CYDO	:		
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
6718_c2_103	152 2072	531	1596 1457	3.5e-149
Protein name			Locus Name sp:PMGI_ECOLI	Acc# P37689
Description				
(EC 5.4.2.1) (PHO	SPHOGLYCEROMUTASE	E) (BPG-IND	EPENDENT PGAM)	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
6837753_f1_23	153 2073	224	675 147	3.2e-08
Protein name			Locus Name	Acc#
capm protein (cap	M1) RP344		pir:B71691	B71691
Description				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
789811_c1_88	154 2074	892	2679 2203	2.3e-256
Protein name			Locus Name	Acc#
			sp:GYRA_ECOLI	P09097
Description				
DNA GYRASE SUBUNI	T A,			
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
986638_c3_5	155 2075	262	789 1149	1.5e-116
Protein name			Locus Name	Acc#
multidrug transpo	rter homolog		pir:G69005	G69005
Description				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12985037_c2_42	156	2076	158	477	354	1.9e-31
<u>Protein name</u>		,			S <u>Name</u> LQ_PSEAE	Acc# P34750
Description						
FIMBRIAL ASSEMBLY P	ROTEIN	PILQ PREC	URSOR			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14301467_c3_49	157	2077	231	696	316	2.9e-28
Protein name					Name	Acc#
carbonic anhydrase				pir:D'	75298	D75298
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
1557_c1_31	158	2078	501	1506	1393	2.1e-142
Protein name					Name_	Acc#
Description				sp:YL	EA_HAEIN	Q57163
HYPOTHETICAL PROTEI	N HI001	9				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
19615687_f1_6	159	2079	87	264		
<u>Protein name</u>				Locus	Name	Acc#
Description						
по-ніт						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23445308_f2_18	160	2080	224	672		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
23859562_c1_32	161	2081	233	702	140	1.7e-	09
Protein name pilus expression production	rotein			Locus gp:PS)	s Name EPONA		<u>Acc#</u> L28837
Pseudomonas syringa (pilN, pilO), pilus pilus expression pro	express	ion prote	ins (pilM	, pilP)ge		-	•
ORF Name	NTID	AAID	NT Length	AA Length	Score	Proba	bility
24040911_c1_33	162	2082	327	984	220	6.3e-	30
Protein name				3,5,	Name LQ_PSEAE		<u>Acc#</u> P34750
Description							
FIMBRIAL ASSEMBLY	PROTEIN	PILO PREC	URSOR				1
		~					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
ORF Name 34510950_c2_39	NTID		NT		<u>Score</u>	Proba	
34510950_c2_39  Protein name	-	AAID	<u>NT</u> Length	Length 1938 Locus	201 S Name		
34510950_c2_39	-	AAID	<u>NT</u> Length	Length 1938	201 S Name		15
34510950_c2_39  Protein name	-	AAID	<u>NT</u> Length	Length 1938 Locus	201 S Name		15 <u>Acc#</u>
Protein name membrane protein	163 ae penic express	AAID  2083  illin bin ion prote	NT Length 645 ding protins (pilm	Length  1938  Locus  gp:PSI  ein (ponz , pilP)ge	201 S Name EPONA	4.9e-	Acc# L28837
Protein name  membrane protein  Description  Pseudomonas syringa (pilN, pilO), pilus	163 ae penic express	AAID  2083  illin bin ion prote	NT Length 645 ding protins (pilm	Length  1938  Locus  gp:PSI  ein (ponz , pilP)ge	201 S Name EPONA	4.9e-	Acc# L28837
Protein name  membrane protein  Description  Pseudomonas syringa (pilN, pilO), pilus pilus expression pro	ne penic express otein (p	AAID  2083  illin bin ion prote ilQ) gene	NT Length 645  ding protins (pilm, partial	Length Locus gp:PSI ein (ponz , pilP)ge cds. AA	201 S Name EPONA A), membrenes, con	4.9e-	Acc# L28837 Steins cds and bility

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4304693_c1_34	165	2085	375	1128 883	2.4e-88
Protein name				Locus Name sp:AROB_NEIGO	Acc# 050468
Description				<del></del>	<del></del>
3-DEHYDROQUINATE S	YNTHASE	,			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4877328_c1_35	166	2086	318	957	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
7042153_c2_43	167	2087	231	696 452	1.1e-42
<u>Protein name</u>				Locus Name sp:AROK_HAEIN	Acc# P43880
Description					
SHIKIMATE KINASE,	(SK)			•	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
7083457_c3_46	168	2088	216	651 154	4.2e-11
Protein name fimbrial assembly	protein	pilO		Locus Name	<u>Acc#</u> S77728
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23703142_c1_3	169	2089	300	900 635	4.5e-62
Protein name				Locus Name sp:YJEK_ECOLI	Acc# P39280
Description					J
HYPOTHETICAL 38.7	KD PROT	EIN IN MO	PA-EFP INT	ERGENIC REGION	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34119052_f1_1	170	2090	204	612	663	4.9e-65
Protein name				Locu	s Name	Acc#
translation elongat	ion fac	tor EF-P		pir:S	34443	G24442 G56
Description						S34443:S56 375:A65225
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
32712915_c2_17	171	2091	77	234		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
33984701_f3_10	172	2092	579	1740	1233	1.9e-125
Protein name					s Name	Acc#
Description				sp:PM	SR_NEIGO	P14930
PEPTIDE METHIONINE	SULFOXI	DE REDUCT	ASE (PEPT	'IDE MET (	O) REDUCT	PASE)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36131500_c3_21	173	2093	308	927	655	3.4e-64
Protein name				Locus	s Name	Acc#
				sp:HT	PX_ECOLI	P23894
Description						
PROBABLE PROTEASE H	TPX, (F	EAT SHOCK	PROTEIN	HTPX)		
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
3907578_c1_15	174	2094	299	900	572	2.1e-55
Protein name				Locus	s Name	Acc#
				sp:DH	PS_ECOLI	
Description				L		P26282:P78
PYROPHOSPHORYLASE)						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4100312_f3_13	175	2095	106	321	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
48828062_c2_16	176	2096	115	348 223	1.2e-16
Protein name probable transglyo Description	cosylase			Locus Name	Acc# T12796:A69
ORF Name  831318_f2_7  Protein name	NTID	<u>AAID</u>	NT Length 472	AA Score Length 1225 Locus Name	Probability  1.4e-124  Acc#
<u>Description</u>	M MELV			sp:HFLX_ECOL1	P25519
GTP-BINDING PROTEI	IN HFLX		NTM.	7.7	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
870250_f2_6	178	2098	255	768 394	1.6e-36
Protein name hypothetical prote	ein in er	idA-gshB	intergenic	Locus Name	Acc# A65080
Description					
ORF Name [10548386_f2_19	NTID	<u>AAID</u>	NT Length	AA Length Score	Probability
Protein name	179		] [/]	Locus Name	Acc#
Description NO-HIT				,	
F					

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
10626558_c2_94	180	2100	182	549 73	0.039
Protein name				Locus Name sp:TEGP_HSV11	Acc# P06481
Description				· · · · · · · · · · · · · · · · · · ·	
TEGUMENT PHOSPHOPRO	TEIN US	9 (10 KD	PROTEIN)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
1178127_f1_10	181	2101	445	1338 1319	1.5e-134
Protein name				Locus Name sp:SYS_HAEIN	Acc# P43833
Description					<del></del>
SERYL-TRNA SYNTHETA	SE, (SE	RINETRN	A LIGASE)	(SERRS)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
12109686_c1_63	182	2102	66	201	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Length Score	Probability
12892086_f2_26	183	2103	81	246	
Protein name				Locus Name	Acc#
Description					
NO-HIT			ų.		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
1369428_c2_97	184	2104	78	237	
Protein name				Locus Name	Acc#
Description					
NO-HIT					1

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
13710925_f3_46	185 2105	148	447 652	7.1e-64
Protein name			Locus Name sp:MTIC_MORBO	Acc# P34721
Description				
METHYLTRANSFERASE N	MBOI C) (M.MBOI C	()		
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
1412642_c1_65	186 2106	147	444 88	0.00042
Protein name			Locus Name sp:YRKI_BACSU	<u>Acc#</u> P54436
Description				<b></b>
HYPOTHETICAL 8.2 KI	PROTEIN IN BLTR	-SPOIIIC	INTERGENIC REGIO	1
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
14250312_c2_100	187 21.07	246	741	
Protein name			Locus Name	Acc#
Description				
NO-HIT				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
1433466_c2_111	188 2108	85	258 141	9.6e-09
<u>Protein name</u>			Locus Name sp:MVIN_ECOLI	Acc# P75932
Description				
VIRULENCE FACTOR M	7IN HOMOLOG			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
14875390_f3_51	189 2109	134	405 302	1.0e-26
Protein name			Locus Name	<u>Acc#</u> P37764
Description			<u> </u>	
HYPOTHETICAL 49.1	CD PROTEIN IN CDS.	A-HLPA IN	TERGENIC REGION	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15020887_c1_83	190	2110	189	570		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15885450_c3_142	191	2111	342	1029	624	6.6e-61
Protein name					s Name IN_HAEIN	Acc# P44958
Description						
VIRULENCE FACTOR M	VIN HOMO	LOG				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
166043_f1_8	192	2112	259	780	416	7.3e-39
Protein name					s Name	Acc#
cytochrome c matura	ation pr	otein B		gp:AF	044582	AF044582
Description						
Shewanella putrefactories of the cytochrome ScyA (scymaturation protein cytochrome c maturate (ccmE)genes, comp.	yA), cyt 3 (ccmB) cionprot	cochrome of cochrone of cochro	c maturati ome c matu	onprotein ration p	n A (ccmA rotein C	(ccmC),
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
17069628_f1_4	193	2113	116	351		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
187703_f2_21	194	2114	113	342	90	0.00026
Protein name					s Name AR_RHISN	Acc# P55365
Description						
HYPOTHETICAL 12.1	KD PROTE	IN Y4AR				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22462757_c1_67	195	2115	67	204	89	0.00033
Protein name hypothetical protein	n SC6E1	.0.02		Locu:	s Name 35489	Acc# T35489
Description						<del></del>
ORF Name 23470003_c1_81	NTID	<u>AAID</u>	NT Length	AA Length 468	Score	Probability 2.3e-31
Protein name		_		Locus	s Name	Acc#
Description	volv vyavy		,		IN_ECOLI	P75932
VIRULENCE FACTOR MY	/IN HOMC	)LOG				
ORF Name [23914017 c2 104	<u>NTID</u>	<u>AAID</u>	NT Length 88	AA Length	Score	Probability [5.5e-09
		] [/				
Protein name hypothetical protei	n vda''			Locus pir:C	S Name	<u>Acc#</u> C69770
	- Juan			] b11:0	09//0	
Description						
ORF Name 24219792_f2_34	<u>NTID</u>	<u>AAID</u>	NT Length 296	AA Length 891	Score 440	Probability 2.1e-41
Protein name		-			s Name SA_PSEAE	Acc# Q59640
Description						
SYNTHASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24244033_c1_70	199	2119	302	909	620	1.8e-60
Protein name				· · · · · · · · · · · · · · · · · · ·	S Name	Acc# P28606
Description						
HYPOTHETICAL 34.1 F	D PROTE	IN IN GLN	A 3'REGIO	N		

ORF Name	NTID AAI	<u>D</u> <u>NT</u> Length	AA Length Score	Probability
24252302_c2_106	200 21	20 493	1482 1229	5.1e-125
Protein name  2-oxoglutarate/mal yflS	late transloc	ator homolog	Locus Name	<u>Acc#</u> F69811
Description			_	
ORF Name 24330005_c3_122	NTID AAI	Length	AA Length Score 945 413	Probability
Protein name		-	Locus Name	Acc# AB017194
Description				
Plectonema boryanı enhancer genes, cor			ptidase, ferredox	in andamidase
ORF Name	NTID AAI	D NT Length	AA Length Score	Probability
24650962_t3_45	202 21	22 261	786 806	3.4e-80
<u>Protein name</u>			Locus Name sp:T2D1_STRPN	Acc# P09356
<u>Description</u>				
(R.DPNI)				
ORF Name	NTID AAI	Length	AA Score Length 54	Probability
24735875_f2_16  Protein name	203 21	23 /3	Locus Name sp:YMT0_YEAST	Acc#
Description				
HYPOTHETICAL 19.2	KD PROTEIN I	N SUB1-ARGR1	INTERGENIC REGION	
ORF Name	NTID AAI	D NT Length	AA Length Score	Probability
25391007_c2_110	204 21	24 216	651 443	1.0e-41
Protein name N-acetyl-anhydromu	ıramyl-L-alan	ine amidase	Locus Name gp:AF082575	Acc# AF082575
Description				
Pseudomonas aerugi transmembrane prote				dase(ampD) and

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
25662782_f2_24	205	2125	258	777 288	2.7e-25
Protein name				Locus Name sp:CCMA_RHOCA	Acc# P29959
Description					
PROTEIN HELA)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
289052_c1_66	206	2126	154	465 220	4.3e-18
Protein name				Locus Name	Acc#
conserved hypothe	cical pro	otein		pir:B75344	B75344
Description		•			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29301457_£3_44	207	2127	93	282	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29507800_c2_95	208	2128	397	1194 883	2.4e-88
Protein name				Locus Name sp:RP32_PSEAE	<u>Acc#</u> P42378
Description					
RNA POLYMERASE SI	GMA-32 FA	ACTOR			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
34569707_c3_131	209	2129	95	288 74	0.023
Protein name				Locus Name	Acc#
F22C12.13				gp:AC007764	AC007764
Description					
Genomic sequence complete sequence.	for Arab	idopsis t	haliana BA	C F22C12 fromchro	omosome I,

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
36335200_t1_13	210	2130	347	1044	506	2.1e-	18
Protein name					Name		Acc# P37764
Description							
HYPOTHETICAL 49.1 K	D PROTEI	N IN CDSA	A-HLPA IN	TERGENIC	REGION		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
36520625_f2_31	211	2131	256	771	723	2.1e-	/1
Protein name				Locus	Name		Acc#
UMP kinase				gp:AB0	010087		AB010087
Description							
Pseudomonas aerugin elongation factor Ts	osa rpsB , UMP ki	, tsf, py nase, rib	osome re	genes for cyclingfa	ribosom actor, co	mplete	ein S2, cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
3907818_f2_32	212	2132	187	564	614	7.6e-6	50
Protein name				Locus	Name		Acc#
ribosome recycling	factor			gp:AB0	010087		AB010087
Description							
Pseudomonas aerugin elongation factor Ts	_						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
391068_f2_33	213	2133	272	819	534	2.3e-5	51
<u>Protein name</u>				Locus	Name		Acc#
				sp:UPI	S_ECOLI		Q47675:P75
Description							247675: <i>1</i> 75 668
(DI-TRANS-POLY-CIS-	DECAPREN	YLCISTRAN	(SFERASE)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
3915930_t3_50	214	2134	204	615	592	1.6e-5	57
Protein name					Name		Acc# P27302
Description							
TRANSKETOLASE 1, (T	K 1)		<u>.</u>				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	<u>Probability</u>
3947932_£3_41	215	2135	299	900 125	2.0e-05
Protein name				Locus Name	Acc# P76370
Description					
HYPOTHETICAL 29.7	KD PROT	EIN IN SB	CB-HISL IN	TERGENIC REGION	PRECURSOR
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4110687_f2_30	216	2136	497	1494 1775	7.1e-183
Protein name				Locus Name sp:TKT1_ECOLI	Acc# P27302
Description					
TRANSKETOLASE 1,	(TK 1)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4345300_f2_20	217	2137	989	2970 2958	0.0
Protein name				Locus Name	Acc#
Description				sp:SYV_HAEIN	P43834
VALYL-TRNA SYNTHE	TASE, (V	ALINETR	NA LIGASE)	(VALRS)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4495268_c1_84	218	2138	110	333 512	4.9e-49
Protein name				Locus Name	Acc#
ferredoxin [3Fe-48] Description	· · · · · · · · · · · · · · · · · · ·			pir:FEAV	A29936:A00 218
ORF Name	NTID	AAID	NT Length	AA Score	Probability
4693768_f1_11	219	2139	435	1308 854	2.8e-85
Protein name				Locus Name sp:DXR_ECOLI	Acc#
Description					P45568:P77 209
REDUCTOISOMERASE)					

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4772325_c1_69	220	2140	93	282 77	0.0071
Protein name  cytochrome b	-			Locus Name gp:ASA228475	Acc# AJ228475
Description					
Andricus solitari	us cytb	gene.			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5109626_f1_6	221	2141	81	246 355	2.1e-32
Protein name				Locus Name sp:MT1A_MORBO	Acc# P34720
Description					
METHYLTRANSFERASE	MBOI A)	(M.MBOI A	۷)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5350281_c3_139	222	2142	76	231	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
6823912_±3_37	223	2143	63	192	
<u>Protein name</u>				Locus Name	Acc#
Description					
NO-HIT	. , -				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
103187_f2_5	224	2144	98	297	
Protein name				Locus Name	Acc#
Description					
NO-HIT	<u>.</u>				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
16917_f2_4	225	2145	164	495 307	2.6e-27
Protein name				Locus Name sp:CYST_ECOL	Acc# P16701
Description					
SULFATE TRANSPORT	SYSTEM	PERMEASE I	PROTEIN CY	ST	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20163930_c1_9	226	2146	271	813 502	5.6e-48
Protein name				Locus Name	Acc# P44922
Description				<u> </u>	<del></del>
ATP-DEPENDENT RNA	HELICASI	E RHLB HON	MOLOG		7
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24257755_c1_8	227	2147	155	468	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16594167_f1_5	228	2148	510	1533	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
22897255_f2_6	229	2149	269	810 305	4.2e-27
Protein name putative acyltran	sferase			Locus Name gp:SCM10	Acc# AL133469
Description				— <u>L</u>	
Streptomyces coel	icolor co	osmid M10.			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24485937_±3_13	230	2150	62	189	147	1.7e-09
Protein name  glutamate dehydroge	enase			_	s Name N010746	Acc# AJ010746
Description				· I		
Antarctic bacterium	n TAD1,	dhe gene.				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2501562_£3_9	231	2151	288	867	547	9.5e-53
Protein name					s <u>Name</u> SH_ECOLI	Acc# P28691
Description				<u> </u>		
CELL DIVISION PROTE	IN FTS	Η,				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25415636_f1_4	232	2152	679	2040	1148	7.4e-181
Protein name					s Name PG ECOLI	Acc# P10413
Description						
PROTEIN C62.5)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26366686_c2_24	233	2153	791	2376	1520	7.5e-156
Protein name					s Name	Acc#
penicillin-binding	proteir	1 1 A		gp:PA	U73780	U73780
Description						
Pseudomonas aerugir cds, and malic enzym				otein 1A	(ponA) c	gene,complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12304661_f2_18	234	2154	584	1755	763	1.2e-75
Protein name					S Name CN_ECOLI	Acc#
Description						P05824:P76 602
DNA REPAIR PROTEIN	RECN (I	RECOMBINAT	ION PROTE	IN N)		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16578133_c3_57	235	2155	65	198	74	0.013
Protein name					s Name BR_TOBAC	<u>Acc#</u> Q40519
Description						•
PHOTOSYSTEM II 10	KD POLY	PEPTIDE 1	PRECURSOR (	PII10)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19564510_f2_17	236	2156	194	585	444	7.8e-42
Protein name N-formylmethionyla	minoacy.	I-tRNA de	eformylase,	Locus pir:S	S Name 23107	Acc#
Description				<b>-</b>		S23107:S41 694:A49696 :B65121
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23554638_f3_29	237	2157	285	858	531	4.7e-51
Protein name	carrie	r protein	synthase		Name 188707	Acc# AF188707
II				] =-		
Description						
Photobacterium pro beta-ketoacyl-acyl aminodeoxychorismat	carrier	protein	synthase I	I (fabF)	-	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23912502_f1_9	238	2158	90	273	200	5.6e-16
Protein name					Name HP_ECOLI	Acc# P37618
Description						
HYPOTHETICAL 9.1 K	D PROTE	IN IN FTS	SY-NIKA INT	ERGENIC P	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23985753_f3_27	239	2159	167	504	273	1.0e-23
Protein name				Locus	Name	Acc#
				gp:EC	J28377	U28377
Description						
Escherichia coli K	-12 geno	ome; appr	coximately	65 to 68	minutes.	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24302263_t1_6	240	2160	193	582	340	8.2e-31
Protein name				Locus	s Name	Acc#
hypothetical protei	n b2948			pir:C	65080	C65080
Description						· <b></b>
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24353458_f2_20	241	2161	308	927	671	6.9e-66
Protein name				Locus	s Name	Acc#
site-specific recom	binase		-	gp:AF	033497	AF033497
Description			-			
Proteus mirabilis s	site-spe	cific rec	ombinase	(xerD) g	ene, comp	oletecds.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
24642562_f2_13	242	2162	102	309		
<u>Protein name</u>				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3007832_f2_19	243	2163	169	510		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36205013_f3_23	244	2164	361	1086	291	1.3e-25
Protein name				Locus	s Name	Acc#
hypothetical protei	.n			pir:G	75388	G75388
Description						_

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3953593_c1_39	245	2165	211	636	386	1.1e-35
Protein name imidazoleglycerol-p	hosphat	e synthase	3	Locus pir:D	S Name 69070	Acc# D69070
Description				,		
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
4883425_f1_2	246	2166	206	621	234	1.4e-19
Protein name					Name	<u>Acc#</u> P36653
Description						
HYPOTHETICAL 21.6 K	D PROTE	IN IN PAR	E-ICC INT	ERGENIC I	REGION (F	193)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6506_f1_3	247	2167	637	1914	2041	4.6e-211
Protein name				Locus	Name	Acc#
topoisomerase IV su	bunit			gp:AB	003429	AB003429
Description						
Pseudomonas aerugin	osa DNA	for topo	isomerase	IV subur	nit, comp	letecds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
805180_c1_38	248	2168	222	669	554	1.7e-53
Protein name					S7_PEA	Acc# Q43072
Description				<u> </u>		<del></del>
IMIDAZOLEGLYCEROL-P	HOSPHAT	E DEHYDRA'	rase, (IG	PD)		·
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
823381_f3_24	249	2169	134	405		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
862761_c1_43	250	2170	72	219		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12281888_c1_40	251	2171	78	237		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1367177_f2_15	252	2172	304	915	674	3.3e-66
Protein name				Locus	s Name	Acc#
					rn_ecori	
Description					LU_ECOLI	
	SE) (URI)	DINE DIPH	OSPHOGLUCO	sp:GA		P25520
Description	SE) (URI) NTID	DINE DIPH	OSPHOGLUCO NT Length	sp:GA		P25520
Description URIDYLYLTRANSFERAS		23 11 12 12 12 12	NT	SP:GA SE PYROP	HOSPHORY	P25520
Description URIDYLYLTRANSFERAS ORF Name	NTID	AAID	<u>NT</u> Length	SP:GA SE PYROPI  AA Length  993  Locus	Score  273  Name	P25520  LASE)  Probability  1.0e-23  Acc#
Description URIDYLYLTRANSFERAS  ORF Name  14463877_t3_23	NTID	AAID	<u>NT</u> Length	SP:GA SE PYROPI  AA Length  993  Locus	Score	P25520  LASE)  Probability  1.0e-23  Acc#
Description URIDYLYLTRANSFERAS  ORF Name  14463877_f3_23  Protein name	<u>NTID</u> 253	AAID 2173	NT Length 330	SP:GA SE PYROPI  AA Length  993  Locus Sp:YJ0	Score  273 S Name  SQ_ECOLI	P25520  LASE)  Probability  1.0e-23  Acc# P39341
Description URIDYLYLTRANSFERAS  ORF Name  14463877_f3_23  Protein name  Description	<u>NTID</u> 253	AAID 2173	NT Length 330	SP:GA SE PYROPI  AA Length  993  Locus Sp:YJ0	Score  273 S Name  SQ_ECOLI	P25520  LASE)  Probability  1.0e-23  Acc# P39341
Description  URIDYLYLTRANSFERAS  ORF Name  14463877_t3_23  Protein name  Description  HYPOTHETICAL 39.8	NTID 253 KD PROT	AAID 2173 EIN IN PE	NT Length 330 PA-GNTV IN	SP:GA  SE PYROPI  AA  Length  993  Locus  SP:YJO  TERGENIC  AA	Score  273  Name  GQ_ECOLI	P25520  LASE)  Probability  1.0e-23  Acc# P39341  (0361)
Description  URIDYLYLTRANSFERAS  ORF Name  14463877_t3_23  Protein name  Description  HYPOTHETICAL 39.8	NTID  253  KD PROT	AAID 2173 EIN IN PE AAID	Length 330  EPA-GNTV IN  NT Length	Sp:GA  SE PYROPI  AA  Length  993  Locus  Sp:YJG  TERGENIC  AA  Length  Length	Score  273  Name  GQ_ECOLI	P25520  LASE)  Probability  1.0e-23  Acc# P39341  (0361)
Description  URIDYLYLTRANSFERAS  ORF Name  14463877_t3_23  Protein name  Description  HYPOTHETICAL 39.8  ORF Name  156515_t2_20	NTID  253  KD PROT	AAID 2173 EIN IN PE AAID	Length 330  EPA-GNTV IN  NT Length	Sp:GA  SE PYROPI  AA  Length  993  Locus  Sp:YJG  TERGENIC  AA  Length  Length	Score  273 S Name  GQ_ECOLI  REGION  Score	P25520  LASE)  Probability  1.0e-23  Acc# P39341  (0361)  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16040927_c2_50	255	2175	112	339	
Protein name				Locus Name	Acc#
Description					
NO-HIT		<del>-</del>			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16610183_c2_54	256	2176	310	933 569	4.4e-55
Protein name				Locus Name sp:TESB_ECOLI	Acc# P23911
Description					
ACYL-COA THIOESTE	RASE II,				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16819827_±1_6	257	2177	137	414	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19531885_c3_57	258	2178	60	183	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19538388_f2_14	259	2179	75	228 73	0.016
Protein name				Locus Name gp:SMI240618	Acc# AJ240618
Description					
Streptococcus mit:	ıs xpt g	ene, strai	in 12261.		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20942936_f3_27	260	2180	376	1131	1060	4.1e-107
Protein name					s Name LE_BACSU	Acc# P55180
<u>Description</u>						
GALACTOSE 4-EPIMERA	SE)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21675051_f3_28	261	2181	321	966	447	3.8e-42
Protein name					s Name FI_ECOLI	Acc# P45803
Description						
HYPOTHETICAL 32.5 K	D PROT	EIN IN MRC	A-PCKA IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23557930_c3_61	262	2182	619	1860	1768	3.9e-182
Protein name					s Name	Acc#
glucosamine synthas	e			gp:AF	032884	AF032884:L
<u>Description</u>						77909
Thiobacillus ferrod (glmU) gene, partial complete cds; and tr	cds;	glucosamin	e synthas	e(glmS) a	and RecG	_
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score 5	Probability
23634680_f2_18	263	2183	423	1272	383	2.3e-35
Protein name					Name	Acc#
putative UDP-glucos	e dehy	irogenase		gp:ALV	V243431	AJ243431
Description						
Acinetobacter lwoff weeF, weeG, weeH, we mip (partial) genes	eI, we	eJ, weeK,	galU, ugd	l, pgi,gal	E, pgm (	(partial) and

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24400250_f3_24	264	2184	860	2583	1162	6.4e-118
Protein name					s Name SB_HAEIN	Acc# P44857
Description						
GLYCEROL-3-PHOSPH	ATE ACYL	TRANSFER	ASE, (GPAT)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24493801_f3_30	265	2185	375	1128	489	1.3e-46
Protein name Faul DNA methyltra	neferace	<u> </u>	<del></del>		S Name 029070	Acc# AF029070
Description				] gb:we	029070	AF023070
Flavobacterium aq	uatile Fa	auI DNA t	methyItrans	ferase (	fauiM) ge	ene, complete
ORF Name	<u>NTID</u>	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26797302_c2_55	266	2186	393	1182	515	2.3e-49
Protein name					s Name IW_ECOLI	Acc# P77562
Description  HYPOTHETICAL 40.4	מאלא און	<del></del>	óma dos a tr	TED CENT	DECTON	
HYPOTHETICAL 40.4	KD PROTI	EIN IN 21			REGION	
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Length	Score	Probability
3317260_f1_5	267	2187	573	1722	1505	2.9e-154
Protein name					s Name	Acc#
putative phosphog.	lucose is	somerase		gp:AL	W243431	AJ243431
Description						
Acinetobacter lwo weeF, weeG, weeH, mip (partial) gene	weeI, we	eJ, weeK	, galV, ugd	, pgi,ga	lE, pgm	partial) and
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3938762_f2_22	268	2188	71	216	71	0.026
Protein name				Locu	s Name	Acc#
transcription reg	ılator ho	omolog yo	ozG	pir:C	69931	C69931
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probal	oility
6729635_c2_46	269	2189	171	516	94	0.0062	2
Protein name				Locu	s Name		Acc#
hypothetical protei	n C45H4	.14		pir:T	32722		T32722
Description						<u> </u>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
976387_£2_19	270	2190	88	267	74	0.0025	
Protein name hypothetical protei Description	n T16L4	.170		<u>Locu</u>	s Name 09929		Acc# T09929
ORF Name  10823462_c1_13  Protein name	<u>NTID</u>	<u>AAID</u>	NT Length	AA Length 204 Locus	Score	Probal	oility Acc#
Description							<del></del>
NO-HIT						······································	
ORF Name 12376535_f2_5	NTID 272	AAID 2192	NT Length 214	AA Length 645	Score 74	Probab 0.0011	
Protein name					S Name U39068		Acc# U39068
Description							
Vibrio cholerae pat	hogenic	ity island	i, partia	l and cor	mplete co	ls.	
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> <u>Length</u>	Score	Probab	oility
22065635_£2_9	273	2193	521	1566	1440	2.2e-1	.47
Protein name sodium/proline symp transporter opuE	orter o	puE:prolir	ie	Locus pir:H	8 Name 69670		<u>Acc#</u> H69670
Description							

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
24228400_c1_20	274	2194	479	1440 1110	2.1e-112
Protein name				Locus Name sp:HEMN_ECOLI	
Description					P32131:P76 772
(COPROPORPHYRINOGEN	NASE) (C	OPROGEN O	XIDASE)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29328457_c1_14	275	2195	98	297 95	7.5e-05
Protein name				Locus Name sp:MINE_ECOLI	Acc# P18198
Description					
CELL DIVISION TOPOL	LOGICAL	SPECIFICI	TY FACTOR		
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
4880303_c1_19	276	2196	193	582 514	3.0e-49
Protein name				Locus Name sp:PTH_HAEIN	Acc# P44682
Description				<u> </u>	
PEPTIDYL-TRNA HYDRO	DLASE, (	(PTH)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
6835875_£2_4	277	2197	60	183	
<u>Protein name</u>				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
900011_c3_28	278	2198	234	705 269	2.7e-23
Protein name probable ribosomal	protein	L25		Locus Name	Acc# H71665
Description	F0			J [pri.n/1665	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
9869006_f1_2	279	2199	72	219	271	1.7e-	23
Protein name  30S subunit ribos	omal prot	ein S21			s Name 014397		<u>Acc#</u> AF014397
Description		-					
Pseudomonas putid protein S21 (rpsU) cds.							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
11885875_c3_76	280	2200	455	1368	1218	7.5e-	124
Protein name					s Name 64 HAEIN		Acc#
Description				L <u>.</u>			P43955:P43 956
HYPOTHETICAL PROT	EIN HIO16	4/165		-			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
12687781_c3_70	281	2201	174	525	512	4.9e-	49
Protein name					s Name 3_HAEIN		<u>Acc#</u> P43814
Description				<u> </u>	-		
TRANSLATION INITI	ATION FAC	TOR IF-3				<u>.</u>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
14093967_c1_49	282	2202	211	636	741	2.6e-	73
Protein name				Locu	s Name		Acc#
<del></del>							
NqrE		·		gp:AF	165980		AF165980
<del></del>				gp:AF	165980		 AF165980

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14540908_c3_77	283	2203	270	813	474	5.2e-45
Protein name					s Name 117331	Acc# AF117331
Description				_		
Vibrio cholerae N10 enzyme complex, com			ating NAL	H-ubiquir	noneoxido	oreductase
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15865712_f2_21	284	2204	189	570	162	6.0e-12
Protein name				Locus gp:EC	Name OUW93	Acc# U14003
Description						
Escherichia coli K	-12 chr	omosomal r	egion fro	m 92.8 to	00.1 mi	inutes.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16460432_c2_65	285	2205	78	237		
Protein name				Locus	s Name	Acc#
Description						,
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
22038177_±3_27	286	2206	557	1674	1801	1.2e-185
<pre>Protein name putative efflux pun</pre>	n comp	nont MtrF			Name	Acc#
	ip compe	Menc Mcr	· · · · · · · · · · · · · · · · · · ·	gp:AF	176821	AF176821
Description						
Neisseria gonorrhoe gene, complete cds.	eae stra	ain EU75 p	utative e	rriux pum	mp compon	lentMtrF (mtrF)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24423260_c1_42	287	2207	437	1314	121	2.5e-05
Protein name				_	Name	Acc#
pr2				gp:MH(	J19289	U19289
Description						
Mycoplasma hyopneum pr1 and pr2 genes, o				_	_	_

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
25392778_f1_1	288	2208	201	606 387	8.6e-36
Protein name  4-hydroxyphenylace	etate 3-r	nonooxyge	nase (EC	Locus Name	<u>Acc#</u> D90737:AB0
Description Escherichia coli	genomic	DNA. (22.	8 - 23.1 π	in).	01340
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
31268837_±3_28	289	2209	412	1239 1836	2.4e-189
Protein name				Locus Name sp:CATA_HAEI	Acc# P44390
Description					
CATALASE,					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
33223291_£2_19	290	2210	71	216	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
33235937_c2_59	291	2211	782	2349 1415	1.0e-144
Protein name				Locus Name	Acc#
Description				sp:VACB_ECOI	P21499:P76 800
VACB PROTEIN					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
33867132_f1_12	292	2212	225	678	
Protein name				Locus Name	Acc#
Description					
NO-HIT					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3399183_c2_61	293	2213	415	1248	1268	3.8e-129
Protein name					s Name 117331	Acc# AF117331
Description				_		<del></del>
Vibrio cholerae N1 enzyme complex, com			cating NAI	H-ubiquii	noneoxido	oreductase
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34000785_c3_73	294	2214	61	186		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34196052_c2_63	295	2215	416	1251	1650	1.2e-169
Protein name				_	Name 117331	Acc# AF117331
Description						
Vibrio cholerae N1 enzyme complex, com			cating NAD	H-ubiquii	noneoxido	preductase
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3939043_c2_58	296	2216	642	1929	2200	6.5e-228
Protein name					Name	Acc# P43014
Description						
(THRRS)			1 330 1 1111			
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
4720967_c2_62	297	2217	227	684	679	9.8e-67
Protein name				<del></del>	Name	Acc#
Description				5p:116	58_HAEIN	P43958:P43 959
HYPOTHETICAL PROTE	IN HIO1	68/169				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
473137_c1_41	298	2218	76	231	
Protein name				Locus Name	Acc#
Description					
NO-HIT		<u></u>			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4801625_f2_20	299	2219	252	759 622	1.1e-60
Protein name				Locus Name sp:HIS4_RHOSH	<u>Acc#</u> P50936
Description					<del></del>
ISOMERASE,					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5882211 ±2 14	300	2220	118	357 [153]	2.7e-10
	J [		السنتا ا		
Protein name	J L			Locus Name	Acc#
Protein name hypothetical prote	J L				Acc# S47051
Protein name	J L			Locus Name	
Protein name hypothetical prote Description ORF Name	J L	AAID	NT Length	Locus Name	
Protein name hypothetical prote Description	in 1		NT	Locus Name  pir:S47051  AA Score	S47051
Protein name hypothetical prote  Description  ORF Name  [682641_c2_55]  Protein name	in 1  NTID  301	AAID 2221	NT Length	Locus Name  pir:S47051  AA Score Length  261  Locus Name	S47051  Probability  2.2e-05  Acc#
Protein name hypothetical prote  Description  ORF Name  [682641_c2_55]  Protein name hypothetical prote	in 1  NTID  301	AAID 2221	NT Length	Locus Name  pir:S47051  AA  Length  261  100	Probability 2.2e-05
Protein name hypothetical prote  Description  ORF Name  [682641_c2_55]  Protein name	in 1  NTID  301	AAID 2221	NT Length	Locus Name  pir:S47051  AA Score Length  261  Locus Name	S47051  Probability  2.2e-05  Acc#
Protein name hypothetical prote  Description  ORF Name  [682641_c2_55]  Protein name hypothetical prote	in 1  NTID  301	AAID 2221	NT Length	Locus Name  pir:S47051  AA Score  Locus Name  pir:G71244  AA Score	S47051  Probability  2.2e-05  Acc#
Protein name hypothetical prote  Description  ORF Name  682641_c2_55  Protein name hypothetical prote  Description	in 1  NTID  301  in PH02:	AAID   2221   17	NT Length   86	Locus Name  pir:S47051  AA Score  Locus Name  pir:G71244  AA Score	Probability  2.2e-05  Acc# G71244
Protein name hypothetical prote  Description  ORF Name  682641_c2_55  Protein name hypothetical prote  Description  ORF Name	in 1  NTID  301  in PH02:	AAID 2221 AAID	NT Length 86 NT Length	Locus Name  pir:S47051  AA Score  Length 100  Locus Name  pir:G71244  AA Score	Probability  2.2e-05  Acc# G71244  Probability
Protein name hypothetical prote  Description  ORF Name  [682641_c2_55]  Protein name hypothetical prote  Description  ORF Name  [14103377_f2_9]	in 1  NTID  301  in PH02:	AAID 2221 AAID	NT Length 86 NT Length	Locus Name  pir:S47051  AA Score  Length  100  Locus Name  pir:G71244  AA Score  Length  501  Locus Name	Probability  2.2e-05  Acc# G71244  Probability  9.0e-41  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16973437_c3_30	303	2223	79	240		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
19745308_±1_3	304	2224	66	201		
Protein name				Locus	<u>Name</u>	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24261257_f3_14	305	2225	123	372	140	8.0e-09
Protein name				Locus	Name	Acc#
				sp:PN0	CB_SALTY	P22253
Description				sp:PN	CB_SALTY	P22253
Description NICOTINATE PHOSPHOL	RIBOSYL	TRANSFERA:	SE, (NAPRT		CB_SALTY	P22253
	RIBOSYL' NTID	TRANSFERAS AAID	SE, (NAPRT <u>NT</u> <u>Length</u>		Score	P22253 Probability
NICOTINATE PHOSPHOI			NT	ASE)		
NICOTINATE PHOSPHOI	NTID	AAID	NT Length	ASE)  AA Length 276  Locus	Score  98  Name	Probability 9.9e-05 Acc#
NICOTINATE PHOSPHOI  ORF Name  25600925_f1_2	NTID	AAID	NT Length	ASE)  AA Length 276  Locus	Score 98	Probability 9.9e-05
NICOTINATE PHOSPHOR  ORF Name  25600925_f1_2  Protein name  Description	NTID 306	AAID	NT Length 91	ASE)  AA Length 276  Locus	Score  98  Name	Probability 9.9e-05 Acc#
NICOTINATE PHOSPHOR  ORF Name  25600925_f1_2  Protein name  Description	NTID 306	<u>AAID</u>	NT Length 91	ASE)  AA Length 276  Locus	Score  98  Name	Probability 9.9e-05 Acc#
NICOTINATE PHOSPHORE  ORF Name  25600925_f1_2  Protein name  Description  (EC 2.4.2) (MONO)	NTID  306  FUNCTION	AAID  2226  NAL TGASE	NT Length 91	ASE)  AA Length 276  Locus Sp:MT0	Score  98  Name  GA_ACICA	Probability  9.9e-05  Acc#  024849
NICOTINATE PHOSPHORE  ORF Name  25600925_f1_2  Protein name  Description  (EC 2.4.2) (MONOR	NTID  306  FUNCTION  NTID	AAID    2226  NAL TGASE	NT Length  NT Length	ASE)  AA Length  Locus Sp:MT0  AA Length  207	Score  98  Name  GA_ACICA	Probability  9.9e-05  Acc#  024849
NICOTINATE PHOSPHORE  ORF Name  25600925_f1_2  Protein name  Description  (EC 2.4.2) (MONO)  ORF Name  30659433_c2_21	NTID  306  FUNCTION  NTID	AAID    2226  NAL TGASE	NT Length  NT Length	ASE)  AA Length  Locus Sp:MT0  AA Length  207	Score  98 S Name  GA_ACICA  Score	Probability  9.9e-05  Acc#  O24849  Probability

ORF Name	NTID	AAID		<u>NT</u> Length	AA Length	Score	Proba	bility
3303178_£1_1	308	2228		179	540	430	2.4e-	40
Protein name  solanesyl diphospha	ate synt	hase				Name 001997		<u>Acc#</u> AB001997
Description								
Rhodobacter capsula	atus DN	A for so	Lane	esyl dip	hosphate	synthase	e,compl	ete cds.
ORF Name	NTID	AAID		<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
35182887_c2_23	309	2229	$\Box$	191	576	684	2.9e-	67
Protein name						s Name YR_HAEIN		<u>Acc#</u> P44529
Description					<u> </u>			
PHOSPHO-HYDROLASE)	(PPASE	)						
ORF Name	NTID	AAID		<u>NT</u> Length	AA Length	Score		bility
36126655_f1_4	310	2230		374	1125	1283	9.7e-	131
Protein name						S Name OC_HAEIN		Acc# P43875
Description								
PHOSPHOLYASE)								
ORF Name	NTID	AAID		<u>NT</u> Length	AA Length	Score	Probal	oility
6834702_f2_11	311	2231		162	489	371	4.3e-	34
Protein name						s Name HJ_HAEIN		<u>Acc#</u> P44609
Description								
HYPOTHETICAL PROTE	IN HI05.	77						
ORF Name	NTID	AAID		<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
882636_c1_15	312	2232		258	777	417	5.7e-3	39
Protein name Tipoate biosynthesi	s prote	ein B				Name 147448		<u>Acc#</u> AF147448
Description	·			<u> </u>				
Pseudomonas aerugin rod-shape-determinin (mltB), rare lipopro lipoate biosynthesis	ng prote otein A	ein (rod/ rlpA),p	A), peni	membran .cillin-	e-bound l binding p	lytictran protein 5	sglyco (dacA	sylase ), and

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
973756_c3_34	313	2233	138	417		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
975055_£2_10	314	2234	745	2238	2349	1.1e-243
Protein name				Locu	s Name	Acc#
polyphosphate kinas	e			gp:AC	RBDOXN	Z46863
Description						
Acinetobacter sp. o	-	obQ, sodM,	lyss, ru	bA, rubB	, estB, d	oxyR,ppk, mtgA,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10673587_f1_4	315	2235	402	1209	1210	5.3e-123
Protein name				Locus	s Name	Acc#
				sp:TY	RB_ECOLI	P04693
Description				•		
AROMATIC-AMINO-ACID	ONIMA C	TRANSFERAS	E, (AROAT	') (ARAT)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14572162_f1_1	316	2236	260	783	586	7.0e-57
Protein name				Locus	s Name	Acc#
				sp:YC	IK_ECOLI	P31808:P77
Description						516 516
(EC 1)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20126386_f2_8	317	2237	198	597	325	3.2e-29
Protein name				Locus	s Name	Acc#
				sp:YT	FL_ECOLI	P39319
Description						
HYPOTHETICAL 49.8 K	D PROTE	IN IN CYS	Q-MSRA IN	TERGENIC	REGION	

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
2609375_c2_26	318 2238	92	279	
Protein name			Locus Name	Acc#
Description				
NO-HIT	-			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
26808317_£2_6	319 2239	232	699 576	8.1e-56
Protein name			Locus Name	Acc#
Description			sp:UBIG_ECOLI	P17993:P76 924
METHYLTRANSFERASE)				
ORF Name	NTID AAID	NT Length	AA Length Score	Probability
34394050_f3_15	320 2240	308	927 889	5.5e-89
Protein name			Locus Name	Acc# P39319
Description				
HYPOTHETICAL 49.8 H	KD PROTEIN IN CYS	Q-MSRA IN	TERGENIC REGION	
ORF Name	NTID AAID	NT Length	AA Length Score	Probability
3911568_f2_7	321 2241	233	702 267	4.5e-23
Protein name			Locus Name	Acc# P40852
Description			_	
PHOSPHOGLYCOLATE PH	HOSPHATASE, CHROM	OSOMAL, (	PGP)	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
4117193_c2_25	322 2242	506	1521 963	7.9e-97
Protein name    leucine aminopeption	lase		Locus Name gp:PPU010261	Acc# AJ010261
Description				
Pseudomonas putida	pepA gene.			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4144818_f3_16	323	2243	362	1089	766	5.9e-76
Protein name				Locu	s Name	Acc#
probable ionictrans	porter			pir:F	70819	F70819
Description				•		<del></del>
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4976550_f1_3	324	2244	312	939	399	4.6e-37
Protein name					s Name	Acc#
Description				sb: AR	HD_ECOLI	P52696:P75 761
HYPOTHETICAL TRANSC	RIPTION	NAL REGULA	ATOR IN MO	DC-BIOA	INTERGENI	C REGION
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1441017_c1_38	325	2245	98	297	124	1.2e-07
						M
Protein name				Locus	s Name	Acc#
Protein name opacity protein opa	51			Locus		
	51					Acc# S36329:S28 628
Opacity protein opa  Description  ORF Name	NTID	AAID	NT Length	Pir:S	36329 Score	S36329:S28 628 Probability
Opacity protein opa  Description  ORF Name  14462827_c3_53		AAID 2246		AA Length 270	36329 Score	S36329:S28 628
Opacity protein opa  Description  ORF Name  14462827_c3_53  Protein name	<u>NTID</u>	-	<u>Length</u>	Pir:S  AA  Length  270  Locus	Score 292 Name	S36329:S28 628 Probability 1.0e-25 Acc#
Opacity protein opa  Description  ORF Name  14462827_c3_53  Protein name  ribosomal protein S	<u>NTID</u>	-	<u>Length</u>	AA Length 270	Score 292 Name	S36329:S28 628 Probability
Opacity protein opa  Description  ORF Name  14462827_c3_53  Protein name	<u>NTID</u>	-	<u>Length</u>	Pir:S  AA  Length  270  Locus	Score 292 Name	S36329:S28 628 Probability 1.0e-25 Acc#
Opacity protein opa  Description  ORF Name  14462827_c3_53  Protein name  ribosomal protein S	<u>NTID</u>	-	<u>Length</u>	Pir:S  AA  Length  270  Locus	Score 292 Name	S36329:S28 628 Probability 1.0e-25 Acc#
Opacity protein opa  Description  ORF Name  14462827_c3_53  Protein name ribosomal protein S  Description	NTID 326	2246	Length  89  NT	AA Length 270 Locus pir:S	Score [292] S Name 38882	S36329:S28 628 Probability 1.0e-25 Acc# S38882
Opacity protein opa  Description  ORF Name  14462827_c3_53  Protein name  ribosomal protein S  Description  ORF Name	NTID 326 15 NTID		Length  89  NT  Length	AA Length  Dir:S  AA Length  AA Length  Locus  Locus  Locus	Score  [292] S Name 38882	\$36329:\$28 628 Probability 1.0e-25 Acc# \$38882 Probability
Opacity protein opa  Description  ORF Name  14462827_c3_53  Protein name  ribosomal protein S  Description  ORF Name  14494026_c2_50	NTID 326 15 NTID		Length  89  NT  Length	AA Length  Dir:S  AA Length  AA Length  Locus  Locus  Locus	Score  [292] S Name  38882  Score  [500] S Name	S36329:S28 628  Probability  1.0e-25  Acc# S38882  Probability  9.1e-48  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14509682_c2_45	328	2248	165	498	230	3.7e-19
Protein name					s Name U39068	Acc# U39068
Description						
Vibrio cholerae pa	thogeni	city islan	d, partia	l and cor	mplete co	ls.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
157765_c2_48	329	2249	96	291	182	4.5e-14
Protein name				Locus	s Name	Acc#
				sp:YR	PM_ACICA	P33989
Description						
HYPOTHETICAL 9.2 K	D PROTE	IN IN RPON	-MURA INT	ERGENIC 1	REGION (C	ORF3)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16510933_c2_52	330	2250	525	1578	545	1.6e-52
Protein name				Locus	s Name	Acc#
Description				sp:FU	MB_ECOLI	P14407:P78
FUMARATE HYDRATASE	CLASS	I, ANAEROB	IC, (FUMA	RASE)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23445931_c2_51	331	2251	454	1365	942	1.3e-94
Protein name					s Name	Acc#
histidinol dehydro	genase			pir:E	70368	E70368
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23650253_c2_49	332	2252	421	1266	1337	1.8e-136
Protein name				Locus	s_Name	Acc#
				sp:MU	RA_ACICA	P33986
Description						
TRANSFERASE) (EPT)						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2381950_c3_58	333	2253	61	186		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT			·			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23867127_c1_41	334	2254	275	828	147	5.2e-10
Protein name					s Name AP_ECOLI	Acc# P45467
Description						
(0191)				,		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
24395637_c1_40	335	2255	149	450	143	6.2e-10
Protein name				-	s Name AN_ECOLI	Acc# P45465
Description						
HYPOTHETICAL 14.8	KD PROTE	EIN IN AGA	AI-MTR INT	ERGENIC	REGION (	0131)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34010260_f1_1	336	2256	119	360	204	2.1e-16
Protein name general stress prot	ein hom	nolog ykzA		<u>Locu</u> pir:F	s <u>Name</u> 69870	<u>Acc#</u> F69870
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5079188_£3_35	337	2257	163	492	461	1.2e-43
Protein name hypothetical protein	n				s Name A224767	Acc# AJ224767
Description				<u> </u>		
Acinetobacter sp. A	ADP1 lor	gene and	ORFs.			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5330087_c3_61	338	2258	370	1113	922	1.7e-92
Protein name			<del></del>		s Name S8_ACEXY	Acc# P45358
Description				<u></u>		
PHOSPHATE TRANSAMII	NASE)	<del></del>				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
954837_c2_44	339	2259	699	2100	2198	1.1e-227
Protein name				Locus	s Name	Acc#
polyribonucleotide	nucleot	idyItrans	ferase	gp:PP	Y18132	Y18132
Description						
Pseudomonas putida	rps0 ar	nd pnp gen	es.			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
969392_f1_13	340	2260	73	222		
Protein name				Locu	s Name	Acc#
<u>Description</u>						
NO-HIT				<del></del>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1070165_c3_42	341	2261	72	219		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT	***	· · · · · · · · · · · · · · · · · · ·		·	<del></del> .	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10993750_£1_2	342	2262	137	414		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20884677_c3_43	343	2263	560	1683	1389	5.7e-142
Protein name probable acyl-CoA d	ehydrog	enase		Locus pir:B	8 Name 75282	<u>Acc#</u> B75282
Description				_ [		
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
24395191_c1_31	344	2264	97	294	71	0.011
Protein name	<u> </u>	Foin or 1	0.36	_	s Name	Acc#
conserved hypotheti	car pro	tern aq_12		pir:F	70406	F70406
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33804680_c2_35	345	2265	796	2391	709	4.4e-72
Protein name				_	s Name	Acc#
site-specific recom	binase			gp:NG	U82253	U82253
Description						
Neisseria gonorrhoe	ae site	-specific	recombin	ase (gcr	gene,co	omplete cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34085165_f3_20	346	2266	496	1491	1327	2.1e-135
Protein name				·	s Name SD_PSEAE	Acc# P26480
Description						
RNA POLYMERASE SIGN	A FACTO	R RPOD (S	IGMA-70)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35823506_c2_32	347	2267	514	1545	1343	4.3e-137
Protein name Butyryl-CoA:Acetate	Coenzy	me A trans	sferase	_	s Name ACTAGEN	Acc# Z69031
Description						
C.thermosaccharolyt	icum ac	tA gene.				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
35939753_c2_34	348	2268	73	222	106	7.2e-05
Protein name probable acyl-CoA of Description	lehydrog	enase		Locus pir:B	s <u>Name</u> 75282	<u>Acc#</u> B75282
ORF Name  3917193_c2_33  Protein name  probable acyl-CoA d  Description	NTID 349 Tehydrog	AAID 2269 enase	NT Length 95	AA Length 288 Locus pir:B	Score  147  s Name  75282	Probability  2.9e-09  Acc#  B75282
ORF Name  3954817_c1_27  Protein name  probable acyl-CoA d	NTID 350 Ehydrog	AAID 2270 enase	NT Length 159	AA Length 480 Locus	386   S Name	Probability  2.9e-35  Acc#  B75282
Description				] [		
ORF Name  5167157_c2_39  Protein name hypothetical protei	NTID 351 n PH180	<u>AAID</u> 2271  1	NT Length	AA Length 486 Locus pir:A	Score [104] S Name 71191	Probability  8.4e-06  Acc# A71191
ORF Name  9923125_c2_40	NTID 352	<u>AAID</u>	<u>NT</u> Length 73	AA Length 222	Score	Probability
Protein name  Description				Locus	s Name	Acc#

ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Score	Probability
10355437_f2_5	353	2273	147	144 159	2.0e-11
Protein name				Locus Name	Acc#
				sp:THID_HAEIN	P44697
Description					
(HMP-P KINASE)			<u>-</u>		
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
23912827_c3_10	354	2274	79	240	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
35267912_c3_11	355	2275	306	921 483	5.8e-46
Protein name				Locus Name	Acc#
				sp:PROC_HAEIN	P43869
<u>Description</u>				<u> </u>	
PYRROLINE-5-CARBOX	YLATE RI	EDUCTASE,	(P5CR) (P	5C REDUCTASE)	
ORF Name	NTID	AAID	NT	AA Score	Probability
			Length	Length ——	
4062840_c2_9	356	2276	191	576 206	1.3e-16
Protein name				Locus Name	Acc#
Description				sp:YGGT_HAEIN	P44097
HYPOTHETICAL PROTE	IN HI103	36			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	<u>Probability</u>
10164063_c2_87	357	2277	242	729 425	8.1e-40
Protein name				Locus Name	Acc#
				sp:YAEB_ECOLI	P28634
Description					<del></del>
HYPOTHETICAL 26.4	KD PROTE	IN IN PRO	s-RCSF IN	TERGENIC REGION	(ORF3)

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14568812_c3_97	358	2278	426	1281 287	5.5e-37
Protein name				Locus Name	Acc#
probable lipD prote	ein			pir:G70634	G70634
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14901512_c3_103	359	2279	156	471 210	4.9e-17
Protein name				Locus Name sp:HIT_BACSU	<u>Acc#</u> 007513
Description					
HIT PROTEIN		<u>.</u>			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
164813_f3_52	360	2280	431	1296 1416	7.8e-145
Protein name				Locus Name	Acc# AB025342
Description					
Moritella marina ge synthesis gene clust		omplete co	ds, simila	r to eicosapentae	enoicacid
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
17068763_f1_16	361	2281	337	1014 1048	7.8e-106
Protein name				Locus Name	Acc#
				sp:HEM2_PSEAE	Q59643
Description					
SYNTHASE) (ALAD) (A	ALADH)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
23444400_c3_92	362	2282	336	1011 1151	9.4e-117
Protein name				Locus Name	Acc# P08577
Description					
HOLLIDAY JUNCTION I	ONA HEL	ICASE RUVE	3 .		

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	oility
23526552_c2_83	363	2283	422	1269	316	2.9e-2	18
Protein name				Locus	s Name		Acc#
conserved hypotheti	cal pro	tein yueF		pir:G	70007		G70007
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	
23595281_f1_17	364	2284	783	2352	2265	8.4e-2	35
Protein name				Locus	s Name		Acc#
hypothetical protei	n b2463	3		pir:F	65021		F65021
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
23828428_f3_57	365	2285	272	819	250	1.6e-4	.0
Protein name				Locus	s Name		Acc#
aldoketoreductase				gp:AF	001865		AF001865
Description							
Leishmania mexicana	amazor	nensis ald	oketoredu	ctase (P'	rR-1) ger	ne,comp.	lete cds.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	oility
24250012_c1_66	366	2286	576	1731	1104	9.0e-1	.12
Protein name				Locus	s Name		Acc#
glycine betaine tra	nsporte	er BetL		gp:AF	102174		AF102174
Description							
Listeria monocytoge cds.	enes gly	ycine beta	ine trans	porter Be	etL (betI	L)gene,	complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
24313512_f2_37	367	2287	121	366		•	
Protein name				Locus	s Name		Acc#
Description							
NO-HIT							

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24317157_f3_55	368	2288	179	540	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
2517175_f1_18	369	2289	78	237	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29376681_f1_1	370	2290	84	255	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
30360452_f1_6	371	2291	80	243	
Protein name				Locus Name	Acc#
Description					
NO-HIT		***		·	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
30662517_c3_106	372	2292	479	1440 514	3.0e-49
Protein name				Locus Name	<u>Acc#</u> I P24180
Description					
ACRIFLAVIN RESIST	ANCE PRO	EIN E PRE	CURSOR (E	NVC PROTEIN)	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31423292_c2_80	373	2293	308	927	327	2.0e-29
Protein name				Locu	s Name	Acc#
hypothetical protei	n Rv024	1c		pir:E	70938	E70938
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31466_f3_54	374	2294	71	216		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT		-				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34110436_f2_30	375	2295	116	351	83	0.030
Protein name microfilarial sheat	h prote	ein SHP3			s Name U54556	Acc# U54556
Description			•	<u> </u>		
Litomosoides sigmoomicrofilarial sheath				_		(shp3a) and
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4147193_f2_29	376	2296	635	1908	1651	3.4e-242
Protein name				Locus	s Name	Acc#
dihydroxy-acid dehy	dratase	2,		pir:D	WECDA	207210 206
<u>Description</u>						A27310:D26 570:S48894 :S30669:F6
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4350088_c3_96	377	2297	458	1377	863	3.1e-86
<u>Protein name</u>					S Name CB1883	Acc# AL022486
Description						
Mycobacterium lepra	e cosmi	d B1883.				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4381318_f3_56	378	2298	250	753 585	9.0e-57
Protein name				Locus Name sp:CCA_ECOLI	Acc# P06961
<u>Description</u>				<del></del>	<del></del>
(TRNA CCA-PYROPHOSI	PHORYLA:	SE) (CCA-2	ADDING ENZ	YME)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
4712537_c1_60	379	2299	117	354	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4769050_c2_79	380	2300	99	300 117	3.5e-07
Protein name hypothetical protein	n APEO	395		Locus Name	Acc# B72732
	n APEO	395			
hypothetical protein Description  ORF Name	NTID	AAID	NT Length	pir:B72732  AA Length Score	
hypothetical protein Description			<del></del>	pir:B72732  AA Score	B72732
hypothetical protein Description  ORF Name	NTID	AAID	<u>Length</u>	pir:B72732  AA Length Score	B72732
hypothetical protein Description  ORF Name  5266540_f1_8	NTID	AAID	<u>Length</u>	pir:B72732  AA Length 660	B72732 Probability
hypothetical protein  Description  ORF Name  5266540_f1_8  Protein name	NTID	AAID	<u>Length</u>	pir:B72732  AA Length 660	B72732 Probability
hypothetical protein  Description  ORF Name  5266540_f1_8  Protein name  Description	NTID	AAID	<u>Length</u>	pir:B72732  AA Length 660	B72732 Probability
hypothetical protein  Description  ORF Name  5266540_f1_8  Protein name  Description  NO-HIT	NTID 381	<u>AAID</u>  2301	Length 219 NT	Pir:B72732  AA Score Locus Name	B72732  Probability  Acc#
hypothetical protein  Description  ORF Name  5266540_f1_8  Protein name  Description  NO-HIT  ORF Name	NTID 381 NTID 382	<u>AAID</u>   2301    AAID   2302	Length 219 NT Length	AA Score Locus Name  AA Length  Locus Name	B72732  Probability  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6697266_c1_62	383	2303	78	237		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6817191_c2_89	384	2304	975	2925	2816	3.5e-293
Protein name					s Name	Acc# P37637
Description				<del></del>		<b>.</b>
HYPOTHETICAL 111.	KD PRO	rein in Hi	DED-GADA I	NTERGENI	C REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
781302_c3_98	385	2305	185	558	541	4.1e-52
Protein name				-	s Name	<u>Acc#</u> P36766
<u>Description</u>						
HYPOXANTHINE PHOS	PHORIBOS	YLTRANSFE	RASE, (HPR	T)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
100305_c3_168	386	2306	251	756	528	9.8e-51
Protein name					s Name HW_ECOLI	Acc# P46852
Description						
HYPOTHETICAL 26.3	KD PROTI	EIN IN GN'	rr-ggt int	ERGENIC	REGION (F	(231)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10604658_f2_36	387	2307	488	1467	705	1.7e-69
Protein name					s Name	Acc#
RdxB				gp:RS	U67862	U67862
Description						
Rhodobacter sphae: genes, partial cds		dxB and re	dxH genes,	complet	e cds, an	d ccoPand rdxI

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
12509836_£2_57	388	2308	137	414	178	1.2e-13	
Protein name hypothetical protei	n R186.	1		Locu:	s <u>Name</u> 24235	<u>Acc#</u> T24235	
Description							
ORF Name	NTID	<u>AAID</u>	NT Length	AA Length	Score	Probability	_
	303	المراقع الم	100				J
Protein name  hypothetical protei	n SPAC	869.06c		Locus pir:T	39117	Acc# T39117	
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
13080050_f1_26	390	2310	66	201	74	0.021	
Protein name					s Name	<u>Acc#</u>	
Description				gp:ST	AF000001	AF000001:	A
Salmonella typhi to Ytl2 homolog (ytl2) (rci) gene, complete	genes,	erase B (t complete	opB), sin cds; pil	gle stra	nd bindir omplete s	ngprotein (ssb), sequence; Rci	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
13723751_c3_176	391	2311	423	1272	1357	1.4e-138	
Protein name FixNd		·		Locu:	s Name FIXND	<u>Acc#</u> Z80339	
Description							
R.leguminosarum fix	Nd and	fixOd gen	es.				]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
140_f1_11	392	2312	144	435			
Protein name				Locus	s Name	Acc#	
Description							
NO-HIT							7

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Se	core	Probability
14650251_c1_125	393	2313	243	732	385	1.4e-35
Protein name				Locus   sp:YGBP		<u>Acc#</u> 005029
Description						<b>_</b>
HYPOTHETICAL PROTEI	N HIO6	72	_			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
156261_f2_53	394	2314	158	477	522	4.3e-50
Protein name				Locus 1		Acc# P44387
Description						
50S RIBOSOMAL PROTE	EIN L13					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
15859456_f3_74	395	2315	96	291	105	6.6e-06
Protein name				Locus 1		Acc#
hypothetical protei	n PH06:	39		pir:H71	108	H71108
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
16803811_f1_13	396	2316	216	651	87	0.040
Protein name				Locus 1		Acc#
somatostatin sst2B	recepto	or	•	gp:RNSS	T2B	X98234
Description		•				
R.norvegicus mRNA i	or som	atostatin	receptor.			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
16853590_c3_164	397	2317	233	702	265	7.3e-23
Protein name				Locus 1		Acc#
Description						P76256:008 476:008477

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
19563312_c2_137	398 2318	96	291 71	0.038
Protein name			Locus Name sp:YYAB_BACSU	Acc# P37523
Description				
HYPOTHETICAL 17.0 F	O PROTEIN IN SPOOL	J-GIDB IN	TERGENIC REGION	
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length <u>Score</u>	Probability
19632661_f3_91	399 2319	134	405	:
Protein name			Locus Name	Acc#
Description				
NO-HIT				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
203577_c1_95	•	751	2256 2566	1.1e-266
Protein name			Locus Name	Acc#
			sp:CLPB_HAEIN	P44403
Description				
CLPB PROTEIN				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
21988931_±3_88		211	636 563	1.9e-54
Protein name			Locus Name	Acc#
			sp:UCRI_CHRVI	031214
Description				
(RIESKE IRON-SULFUR	R PROTEIN) (RISP)			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
22066661_f2_40	402 2322	191	576 364	2.4e-33
Protein name			Locus Name	Acc#
			sp:YAJQ_HAEIN	P44096
Description				
HYPOTHETICAL PROTEI	N HI1034			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23525307_c2_146	403	2323	213	642	554	3.5e-58
Protein name cytochrome-c oxidas	se. type	e cbb3 cha	in fixO	Locu pir:S	s Name	<u>Acc#</u> S77596
<u> </u>				J [P11.5		
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23720002_c2_140	404	2324	61	186		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT					<del></del>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23860681_f2_39	405	2325	455	1368	1917	6.4e-198
Protein name		<u> </u>		Locu	s Name	Acc#
				sp:AS	SY_HAEIN	P44315
Description				<u> </u>		
LIGASE)		., ,				1
			NT	AA		
ORF Name	NTID	AAID	Length	<u>Length</u>	Score	Probability
23864180_f1_18	406	2326	271	816	254	1.1e-21
Protein name				Locus	s Name	Acc#
CorE				gp:AF	130857	AF130857
Description						
Salmonella typhimu:	rium col	balt resis	tance loc	us, part	ial seque	nce.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23947151_£1_19	407	2327	102	309	119	2.2e-07
Protein name				Locus	s Name	Acc#
unknown				gp:AF	147448	AF147448
Description					-	
Pseudomonas aerugir rod-shape-determinir (mltB), rare lipopro lipoate biosynthesis	ng prote otein A	ein (rodA) (rlpA),pe	, membran nicillin-	e-bound i binding p	lytictran protein 5	sglycosylase (dacA), and

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24083208_f3_82	408	2328	71	216	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24271875_c1_122	409	2329	558	1677 1857	1.5e-191
Protein name				Locus Name sp:PYRG_HAEIN	Acc# P44341
Description					
CTP SYNTHASE, (UTP	AMMON	IA LIGASE)	(CTP SYN	THETASE)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
24337827_f1_15	410	2330	355	1068 1038	8.9e-105
Protein name		<del></del>		Locus Name	Acc#
dihydroorotase,				pir:T10453	T10453
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24344138_f3_68	411	2331	70	213	
Protein name				Locus Name	Acc#
Description					
NO-HIT		·			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24417875_c1_124	412	2332	141	135	4.3e-09
Protein name				Locus Name	Acc#
				sp:YGBQ_HAEIN	P44035
Description					
HYPOTHETICAL PROTE	IN HI06	73			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24500286_c3_163	413	2333	519	1560 1458	2.6e-152
Protein name				Locus Name sp:SR54_ECOLI	Acc# P07019
<u>Description</u>				<u>.                                    </u>	
SIGNAL RECOGNITION	N PARTIC	LE PROTEIN	(FIFTY-F	OUR HOMOLOG) (P48	3)
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Score	Probability
24648402_f1_22	414	2334	1298	3897 386	5.4e-59
Protein name				Locus Name	Acc#
probable exonuclea	ise,	· · <u> </u>		pir:T03465	T03465
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24844562_c3_167	415	2335	548	1647 1470	1.5e-150
Protein name				Locus Name	Acc#
probable pitB prot	ein			pir:E70731	E70731
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29880042_f3_83	416	2336	485	1458 527	3.1e-61
Protein name				Locus Name	Acc#
				sp:SBCD_ECOLI	P13457
Description					
EXONUCLEASE SBCD					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3166026_f3_87	417	2337	67	204	
Protein name				Locus Name	Acc#
Description					
NO-HIT			-		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33204808_c1_101	418	2338	350	1053	231	2.8e-18
Protein name hypothetical prote	in RP372	2		Locus pir:E	8 Name 71694	Acc# E71694
Description				· · ·		
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
3367635_f2_55	419	2339	421	1266	1355	2.3e-138
Protein name					s Name B_CHRVI	Acc# 031215
Description						
CYTOCHROME B						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
33707182_f1_27	420	2340	252	759	364	2.7e-46
Protein name					s Name 1_CHRVI	Acc# 031216
Description						
CYTOCHROME C1 PREC	URSOR					
ORF Name [33875885_c3_157	NTID	<u>AAID</u>	NT Length	AA Length 207	Score	Probability
Protein name	J L			Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
34064681_c1_119	422	2342	87	264	71	0.026
Protein name				_	s Name	Acc#
cb-type cytochrome	c oxida	se CcoQ s	ubunit	gp:AB	024290	AB024290
Description						
Magnetospirillum m	_		ON, CCOO,	ccoQ, co	coP gene	forcb-type

ORF Name	NTID AAID NT Length	AA Score Probability
34120251_c1_105	423 2343 322	969 647 2.4e-63
Protein name		Locus Name Acc#  Sp:UBIA_ECOLI P26601
Description		
POLYPRENYLTRANSFERA	ASE)	
ORF Name	NTID AAID NT Length	AA Score Probability
36379680_c2_127	2344 60	183
Protein name		Locus Name Acc#
Description		
NO-HIT		
ORF Name	NTID AAID NT Length	AA Score Probability
3906686_c3_155	425 2345 653	1962 2231 3.4e-231
Protein name		Locus Name Acc#
	·	sp:GIDA_PSEPU P25756
Description		
GLUCOSE INHIBITED I	DIVISION PROTEIN A	
ORF Name	NTID AAID NT Length	AA Score Probability
3932753_c2_149	426 2346 767	2304 235 1.3e-16
Protein name		Locus Name Acc#
		sp:REC2_HAEIN P44408
Description		
RECOMBINATION PROTE	EIN 2	
ORF Name	NTID AAID NT Length	AA Score Probability
3942318_f2_54	427 2347 131	396 507 1.7e-48
Protein name		Locus Name Acc#  Sp:RS9 HAESO P31782
Description		5P.103_1M200 F31702
30S RIBOSOMAL PROTE	IN 89	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probab	oility
3947193_f2_56	428	2348	132	399	311	9.7e-2	28
Protein name				Locus N			<u>Acc#</u> P45206
Description							
STRINGENT STARVATIO	N PROTI	EIN B HOMO	LOG				
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Length	ore	Probab	oility
4119075_c1_103	429	2349	281	846	464	6.0e-4	4
Protein name				Locus N			Acc#
Description							P31054:P39 203
(EC 2.7.1.66)							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Sc	ore	Probal	oility
4334463_c3_172	430	2350	169	510	78	2.8e-0	)5
Protein name				Locus N			Acc#
unknown				gp:AF083	3916		AF083916
Description							
Rhizobium etli Fnr- complete cds; and un		_	onal regu	lator FnrNo	(fnrN	c) gene	,
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probab	oility
4798193_c3_178	431	2351	358	1077	368	5.6e-4	.8
Protein name				Locus N	<u>lame</u>		Acc#
cytochrome-c oxidas cytochrome-c oxidase b410:fixP protein				pir:D474	. 168		D47468
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Sc	ore	Probab	oility
500017_f3_73	432	2352	241	726	550	4.6e-5	3
Protein name				Locus N sp:RNT V			Acc# P46232
Description				<u> </u>			
RIBONUCLEASE T, (EX	ORIBON	UCLEASE T)	(RNASE T	)			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	bility
520003_c1_126	433	2353	67	204			
Protein name				Locu	s Name		Acc#
Description							
NO-HIT		······································					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
5203453_c3_181	434	2354	445	1338	1467	3.1e-1	150
Protein name					s Name O_ECOLI	1	Acc# P08324
Description				<u> </u>			
GLYCERATE HYDRO-LY	ASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
5281318_c3_180	435	2355	290	873	1037	1.1e-1	104
Protein name				Locu	s Name		Acc#
2-dehydro-3-deoxyp	hosphood	ctonate a	ldolase	gp:AF	098791		AF098791
Description							
Pseudomonas aerugi complete cds.	nosa 2-0	dehydro-3	-deoxyphos	phoocton	ate aldol	ase (kd	sA) gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
5901067_c1_104	436	2356	274	825	202	3.5e-1	16
Protein name					s Name		Acc#
Description				sp:in	IQ_HAEIN		P44901
HYPOTHETICAL PROTE	IN HIO84	19	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			····	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
7054650_c1_118	437	2357	62	189	53	0.015	
Protein name	<b>_</b>			Locus	s Name		Acc#
ORF-D					010KLS		D11109
Description							
E. coli gene for 1	OK-L and	1 10K-S p	rotein.		<u> </u>		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
957705_c1_113	438	2358	335	1008	407	6.5e-38
Protein name  putative regulatory	/ protei	n		_	s Name 087482	Acc# AF087482
<u>Description</u>				Y 0 F	a maguila	Formyotoin
Pseudomonas aerugii (clcR), chlorocatecl (clcB), dienelactone transposase (tnpA), (ohbR),o-halobenzoa alpha subunit (ohbB)	nol-1,2- e hydrol ATP-bind te dioxy	-dioxygen lase(clcD ling prot ygenase r	ase (clcA) ), maleyla ein (tnpB) eductase (	,chlorom cetate re , putation ohbA), o	uconate deductase ve regula -halobenz subunit (	ycloisomerase (clcE), tory protein oatedioxygenase ohbC),
ORF Name	NTID	AAID	<u>Length</u>	<u>Length</u>	Score	Probability
9960917_f3_90	439	2359	223	672	354	2.7e-32
Protein name					s Name PA_ECOLI	Acc# P05838
Description						
STRINGENT STARVATIO	ON PROTE	EIN A				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10632090_f1_17	440	2360	506	1521	981	9.8e-99
Protein name		,			s Name ON_ECOLI	Acc#
Description				<u> </u>		P33608:P78 281
OXIDOREDUCTASE CHA	IN 14)	(NUO14)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1069465_c1_85	441	2361	75	228		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10734830_c1_89	442	2362	60	183		
Protein name	•			Locu	s Name	Acc#
Description						
NO-HTT						

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
1385390_f3_54	443 2363	216	651 409	4.0e-38
Protein name			Locus Name sp:NUOJ_ECOLI	Acc#
<u>Description</u>				P33605:P78 236
OXIDOREDUCTASE CHA	IN 10) (NUO10)			
ORF Name	NTID AAID	NT Length	AA Length Score	Probability
13863425_f2_23	2364	276	831 480	1.2e-45
Protein name			Locus Name	Acc#
hypothetical prote	in RP682		pir:E71674	E71674
Description				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
14454827_f2_28	2365	211	561	3.1e-54
Protein name			Locus Name	Acc#
pyridoxamine 5-phos	sphate oxidase		pir:B75513	B75513
Description				
ORF Name	NTID AAID	NT Length	AA Length Score	Probability
14475702_c1_90	2366	259	780 91	0.00081
Protein name			Locus Name	Acc#
ORF8			gp:D78257	D78257
Description				
Enterococcus faeca ORF6, ORF7, ORF8, O				3,ORF4, ORF5,
ORF Name	NTID AAID	<u>NT</u> <u>Length</u>	AA Length Score	Probability
14578202_f1_12	2367	182	763	1.2e-75
Protein name			Locus Name	Acc#
Description			sp:NUOI_ECOLI	P33604:P76 488:P78183
OXIDOREDUCTASE CHA	IN 9) (NUO9)			133.173183

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15625443_c1_84	448	2368	61	186		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
175760_f3_46	449	2369	215	648	352	4.4e-32
Protein name				Locu	s Name	Acc#
NADH dehydrogenas	e chain A	A		gp:AF	057063	AF057063
Description						
Erwinia carotovor partial cds; HexA dehydrogenase chai C (nuoC) gene, par	(hexA), l n B (nuol	NADH dehy B) genes,	drogenase	chain A(	nuoA), ar	nd NADH
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19806577_£2_27	450	2370	452	1359	1177	1.7e-119
Protein name					s Name SA_HAEIN	Acc# P45164
Description						
MRSA PROTEIN HOMO	LOG					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2110657_f1_3	451	2371	328	987	760	2.6e-75
Protein name					s Name 26_SYNY3	<u>Acc#</u> P72872
Description						
HYPOTHETICAL 37.9	KD PROT	EIN SLLOS	26			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22402252_f2_25	452	2372	63	192		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
23683215_f2_38	453	2373	579	1740	1583	1.6e-162
Protein name				Locu	s Name	Acc#
				`	OM ECOLI	
Description				L		P31978:P78
<u>Descripcion</u>						248
OXIDOREDUCTASE CHA	IN 13)	(NUO13)				
ORF Name	NTID	AAID	NT	<u>AA</u>	Score	Probability
ORF Name	NIID	AAID	Length	Length	SCOLE	FIODADILICY
24225213_f3_50	454	2374	266	801	1183	3.8e-120
Protein name				Locus	s Name	Acc#
Tou2					058689	AF058689
Description				J <u>5-</u>		
Neisseria meningit	idis st	rain Z2491	, genomic	sequence	е.	
ORF Name	NTID	AAID	$\underline{ ext{NT}}$	<u> AA</u>	Score	Probability
	MIID	MID	<u>Length</u>	<u>Length</u>	bcore	FIODADIIICY
24226502_c3_132	455	2375	270	813	888	7.0e-89
Protein name				Locus	s Name	Acc#
					72 HAEIN	P44758
Description						
Description						
HYPOTHETICAL PROTE	IN HI05	72				
ORF Name	NTID	AAID	NT	<u>AA</u>	Score	Probability
ORF Name	NIID	AAID	Length	Length	SCOLE	Probability
24391557_f1_10	456	2376	1046	3141	1655	8.6e-252
Protein name				Locus	s Name	Acc#
NADH dehydrogenase	(ubiqui	none), I	chain			·
G:nuoK protein	_					A65000:S65
Description				J		638:S38316
						:S37064
			NT	AA		
ORF Name	NTID	AAID	Length	Length	Score	<u>Probability</u>
24642893 f1 15	457	2377	619	1860	1809	1.8e-186
	J L					
<u>Protein name</u>					Name	Acc#
				sp:NU	Or_ECOLI	D22607 D22
Description				·		P33607:P78
						234
OXIDOREDUCTASE CHA	IN 12)	(NUO12)	····			

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probab	ility
2507286_£2_22	458	2378	213	642	770	2.2e-7	6
Protein name  outer membrane prot	ein Bl				s Name 045251	<del></del>	Acc# AF045251
Description				<u> </u>	-		
Moraxella catarrhal	is outer	r membrane	e protein	B1 gene	, complet	e cds.	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	ility
25392135_f2_26	459	2379	61	186			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
25579763_f3_61	460	2380	281	846	374	2.3e-3	9
Protein name					s Name NR ECOLI	<del></del> 1	Acc#
Description				Sp.11			P28861:P11 007
(FLXR) (FLDR) (METH	AT AIOTO	GEN RESIS	STANCE PR	OTEIN A)	(DA1)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
26228401_c2_105	461	2381	156	471	123	8.1e-0	8
Protein name				Locus	s Name		Acc#
hypothetical protein	n APE141	.3		pir:D	72619		D72619
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
29688176_f1_1	462	2382	70	213	304	4.8e-2	6
Protein name					s Name		Acc#
transferrin-binding	protein	2 precur	rsor	gp:AF	105251		AF105251
Description							
Moraxella catarrhal partial cds.	is trans	ferrin-b	inding pr	otein 2 p	precursor	(ompB1)	gene,

ORF Name	NTID AAID	<u>NT</u> Length	Length Score	Probability
30082693_f3_51	463 2383	491	1476 1378	8.3e-141
Protein name			Locus Name	Acc#
Description			sp:NUOF_ECOLI	P31979:P78 239
OXIDOREDUCTASE C	HAIN 6) (NUO6)			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
30252036_c2_98	464 2384	63	192	
Protein name			Locus Name	Acc#
Description				
NO-HIT				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
31283452_f1_11	465 2385	342	1029 1126	4.2e-114
Protein name			Locus Name	Acc#
Description			sp:NUOH_ECOLI	P33603:P78
OXIDOREDUCTASE C	HAIN 8) (NUO8)			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
3182067_c3_131	466 2386	516	1551 1966	4.1e-203
Protein name			Locus Name	Acc#
			sp:SYR_HAEIN	P43832
Description			**************************************	
ARGINYL-TRNA SYN	THETASE, (ARGININE-			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
33723387_f1_6	2387	235	708 799	1.9e-79
Protein name			Locus Name	Acc#
Description			sp:NUOB_ECOLI	P33598:P78
OXIDOREDUCTASE C	HAIN 2) (NUO2)			

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33772186_£3_41	2388	416	1251	1601	1.9e-164
Protein name transferrin binding Description	protein B		Locus gp:AF(	Name 039313	Acc# AF039313
Moraxella catarrhal complete cds.	is strain LES-1 t	ransterr	in bindir	ng protei	n B(tbpB) gene,
ORF Name	NTID AAID	<u>NT</u> Length	AA Length	Score	Probability
34176950_f3_42	2389	548	1647	331	2.2e-29
Protein name				Name OMETJA	<u>Acc#</u> Q57634
<u>Description</u>					
HYPOTHETICAL PROTEI	N MJ0170		<del>-</del>		
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34414552_f3_47	470 2390	584	1755	2190	7.5e-227
Protein name			Locus	Name	Acc#
NADH dehydrogenase C-D	(ubiquinone), I,	chain	pir:D6	55000	D65000:S38
<u>Description</u>					313:S38312 :S65634:S6
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35166075_£1_4	471 2391	294	885	315	3.7e-28
Protein name			Locus	Name	Acc#
periplasmic chaperon	ne protein		gp:AF0	95845	AF095845
Description					
Pseudomonas syringa periplasmic chaperon			_		tsK) and
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36144687_f3_49	472 2392	62	189	240	3.2e-20
Protein name				Name D SALTY	Acc# P33902
Description				_	
OXIDOREDUCTASE CHAI	N 4) (NUO4) (FRAG	MENT)			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3915693_c1_80	473	2393	416	1251	211	1.7e-14
Protein name				Locus gp:ECF	Name MC7A	Acc# X57583
Description						
E.coli Plasmid pMo	CCC7 mcc.	A,B,C,D,E,	F genes.			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4740902_c2_127	474	2394	313	942	114	0.00043
Protein name				Locus sp:PRX	Name H_BPMD2	<u>Acc#</u> 064252
Description						·
PUTATIVE NON-HEME	HALOPER	OXIDASE,				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4796875_f1_5	475	2395	78	237	144	4.8e-10
Protein name				Locus	Name	Acc#
conserved hypothet	ical pro	otein		pir:H7	5273	H75273
_ , , ,						
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
	<u>NTID</u>	<u>AAID</u>			Score 320	Probability
ORF Name			Length	Length	320	
ORF Name 5097886_f1_14			Length	Length 438 Locus	320	1.1e-28
ORF Name  5097886_f1_14  Protein name	476		Length	Length 438 Locus	320 Name	Acc# P33606:P76
ORF Name 5097886_f1_14 Protein name Description	476	2396	Length	Length  438  Locus  sp:NUO	320 Name	Acc# P33606:P76
ORF Name  5097886_f1_14  Protein name  Description  OXIDOREDUCTASE CHA	476	2396 (NUO11)	Length  145  NT	Length  438  Locus  sp:NUO	320 Name K_ECOLI	1.1e-28 <u>Acc#</u> P33606:P76 487:P78182
ORF Name  5097886_f1_14  Protein name  Description  OXIDOREDUCTASE CHA	476 AIN 11) NTID	2396 (NUO11) AAID	Length  145  NT Length	Length  438  Locus  sp:NUO  AA  Length  525  Locus	Name K_ECOLI  Score	Acc# P33606:P76 487:P78182  Probability
ORF Name  5097886_f1_14  Protein name  Description  OXIDOREDUCTASE CHA ORF Name  7226452_f1_9	476 AIN 11) NTID	2396 (NUO11) AAID	Length  145  NT Length	Length  438  Locus  sp:NUO  AA  Length  525  Locus	Name K_ECOLI  Score 470  Name	Probability  Probability  Acc#  Acc#  Acc#  Acc#  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
10181576_f2_42	478	2398	101	306	
Protein name				Locus Name	Acc#
Description					
NO-HIT			, <del>, , , , , , , , , , , , , , , , , , </del>		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
10751312_t1_7	479	2399	939	2820 710	2.9e-114
Protein name				Locus Name	Acc#
Description				sp:YCBY_HAEIN	P44524:P43
HYPOTHETICAL PROTE	IN HIO1	16/115			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
10975302_c1_93	480	2400	293	882 185	2.5e-13
Protein name				Locus Name	Acc#
probable D,D-carbo	xypeption	dase		pir:B71353	B71353
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19587762_c1_77	481	2401	89	270	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19735877_t2_34	482	2402	63	192	
Protein name				Locus Name	Acc#
Description					
NO-HIT					

ORF Name	NTID AAID NT Length	AA Length Score Probability
21491075_c3_127	483 2403 517	1554 309 1.4e-41
Protein name CjaB protein		<u>Locus Name</u> <u>Acc#</u> gp:CJE17971 Y17971
Description		
Campylobacter jeju	ni cjaB gene.	
ORF Name	NTID AAID NT Length	AA Length Score Probability
21520276_c3_136	2404 275	828
<u>Protein name</u>		Locus Name Acc#
Description	•	
NO-HIT		
ORF Name	NTID AAID NT Length	AA Score Probability
21603403_c3_126	485 2405 543	1632 857 1.3e-85
Protein name		Locus Name Acc#
		sp:YMDC_ECOLI P75919
Description		
HYPOTHETICAL 55.9	KD PROTEIN IN CSGC-MDOG I	NTERGENIC REGION
ORF Name	NTID AAID NT Length	AA Score Probability
21679677_f3_58	486 2406 476	1431   1649   1.6e-169
Protein name		Locus Name Acc#
		sp:GLNA_AZOVI P22248
Description		<del> </del>
GLUTAMINE SYNTHETAS	SE, (GLUTAMATEAMMONIA L	IGASE)
ORF Name	NTID AAID NT Length	AA Score Probability
22306532_c3_134	487 2407 255	768 426 6.3e-40
Protein name		Locus Name Acc#
		sp:LPSA_PASHA Q05770
Description		
LPSA PROTEIN		

ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
22442010_f1_1	488	2408	354	1065	450	1.8e-42
Protein name unknown					Name 116284	Acc# AF116284
Description						
Pseudomonas aerugir genes.	nosa Dna	aJ-like pi	rotein gen	e, comple	ete cds;	andunknown
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2375337_f3_49	489	2409	60	183		•
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23944431 c2 116	490	2410	80	243	106	5.1e-06
23711101_02_110	J L					
Protein name	l L	_] [	] []	Locus	S Name	Acc#
	l L	_] [			S Name	Acc# H72754
Protein name	l L	_] [		Locus	S Name	<del></del>
Protein name hypothetical protei  Description  ORF Name	n APEOC	_] [	NT Length	Locus pir:H	S Name 72754 Score	H72754
Protein name hypothetical protein Description	n APEO	029	NT	Locus pir:H	S Name 72754	H72754
Protein name  hypothetical protein  Description  ORF Name  23945931_t3_55  Protein name	n APEOC	AAID 2411	NT Length	Locus pir:H  AA Length 1041 Locus	Score  136  Name	H72754  Probability  1.2e-06  Acc#
Protein name hypothetical protein Description  ORF Name  23945931_f3_55	n APEOC	AAID 2411	NT Length	Locus pir:H	Score  136  Name	H72754  Probability  1.2e-06
Protein name  hypothetical protein  Description  ORF Name  23945931_t3_55  Protein name	n APEOC	AAID 2411	NT Length	Locus pir:H  AA Length 1041 Locus	Score  136  Name	H72754  Probability  1.2e-06  Acc#
Protein name  hypothetical protein  ORF Name  23945931_f3_55  Protein name  hypothetical protein	n APEOC	AAID 2411	NT Length	Locus pir:H  AA Length 1041 Locus	Score  136  Name	H72754  Probability  1.2e-06  Acc#
Protein name hypothetical protein  ORF Name  23945931_f3_55  Protein name hypothetical protein  Description	n APEOC	AAID 2411	NT Length 346	Locus pir:H  AA Length 1041 Locus pir:S	Score  136 S Name 75877	H72754  Probability  1.2e-06  Acc#  S75877
Protein name hypothetical protein  ORF Name  23945931_f3_55  Protein name hypothetical protein  Description  ORF Name	n APEOC	AAID 2411 AAID AAID	NT Length 346 NT Length	Locus  AA  Length  Locus  pir:S'  AA  Length  Locus  Locus  Length  Locus	Score  136 S Name 75877  Score  2745 S Name	Probability  1.2e-06  Acc# S75877  Probability  1.2e-285  Acc#
Protein name hypothetical protein  ORF Name  23945931_f3_55  Protein name hypothetical protein  ORF Name  23954511_f1_6	n APEOC	AAID 2411 AAID AAID	NT Length 346 NT Length	Locus  AA  Length  Locus  pir:S'  AA  Length  Locus  Locus  Length  Locus	Score  136 S Name 75877  Score  2745	Probability  1.2e-06  Acc# S75877  Probability  1.2e-285

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23989752_c1_84	493	2413	166	501 288	1.0e-42
Protein name				Locus Name sp:3DHQ_NEUCR	Acc# P05195
Description					
DEHYDRATASE)		·			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24306512_c2_99	494	2414	202	609 509	1.0e-48
Protein name				Locus Name sp:GCH1_OSTOS	<u>Acc#</u> 061573
Description					
GTP CYCLOHYDROLAS	E I, (GT	P-CH-I)			
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
24337752_f2_32	495	2415	378	1137 884	1.9e-88
Protein name				Locus Name	Acc#
Protein name  Description				Locus Name sp:YDAO_ECOLI	Acc# P76055:Q47 558
	KD PROT	EIN IN DB	PA-INTR IN	sp:YDAO_ECOLI	P76055:Q47
Description	KD PROT	EIN IN DB	PA-INTR IN NT Length	sp:YDAO_ECOLI	P76055:Q47
Description HYPOTHETICAL 35.6			NT	sp:YDAO_ECOLI  TERGENIC REGION  AA Score	P76055:Q47 558
Description  HYPOTHETICAL 35.6  ORF Name	NTID	AAID	NT Length	sp:YDAO_ECOLI  TERGENIC REGION  AA Length Score	P76055:Q47 558
Description  HYPOTHETICAL 35.6  ORF Name  24646887_t1_16	NTID	AAID	NT Length	Sp:YDAO_ECOLI  TERGENIC REGION  AA Length  510	P76055:Q47 558 Probability
Description  HYPOTHETICAL 35.6  ORF Name  24646887_f1_16  Protein name	NTID	AAID	NT Length	Sp:YDAO_ECOLI  TERGENIC REGION  AA Length  510	P76055:Q47 558 Probability
Description  HYPOTHETICAL 35.6  ORF Name  24646887_f1_16  Protein name  Description	NTID	AAID	NT Length	Sp:YDAO_ECOLI  TERGENIC REGION  AA Length  510	P76055:Q47 558 Probability
Description  HYPOTHETICAL 35.6  ORF Name  24646887_t1_16  Protein name  Description  NO-HIT	NTID 496	<u>AAID</u> 2416	NT Length 169	Sp:YDAO_ECOLI  TERGENIC REGION  AA Score Locus Name	P76055:Q47 558 Probability  Acc#
Description  HYPOTHETICAL 35.6  ORF Name  24646887_f1_16  Protein name  Description  NO-HIT  ORF Name	NTID 496 NTID	AAID  AAID	NT Length 169 NT Length	Sp:YDAO_ECOLI  TERGENIC REGION  AA Score  510  Locus Name  AA Score	P76055:Q47 558 Probability  Acc#
Description  HYPOTHETICAL 35.6  ORF Name  24646887_t1_16  Protein name  Description  NO-HIT  ORF Name  24881717_t2_39	NTID 496 NTID	AAID  AAID	NT Length 169 NT Length	Sp:YDAO_ECOLI  TERGENIC REGION  AA Score  510  Locus Name  AA Score  Length Score  AA Score	P76055:Q47 558  Probability  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25595262_f3_68	498	2418	168	507		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT					<del></del>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26354750_£3_50	499	2419	60	183		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29332503_f3_66	500	2420	301	906	797	3.1e-79
Protein name					s Name	Acc#
enoyl-(acyl-carrier	protei	n) reduct	ase	gp:AF	104262	AF104262
Description						
Pseudomonas aerugir complete cds.	osa end	yl-(acyl-	carrier p	rotein)	reductase	e(fabI) gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29335786_f3_46	501	2421	249	750	428	3.9e-40
Protein name				_	s Name	Acc#
unknown				gp:AF	116284	AF116284
Description						
Pseudomonas aerugir genes.	nosa Dna	J-like pr	otein gen	e, comple	ete cds;	andunknown
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
29382075_f1_4	502	2422	312	939	429	3.0e-40
Protein name				Locus	s Name	Acc#
probable membrane p	rotein	b1520		pir:C	64906	C64906
Description						<u>-</u>

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31425825_f1_22	503	2423	220	663	771	1.7e-76
Protein name					s Name E_HAEIN	Acc# P44756
Description						
EPIMERASE) (PPE) (	R5P3E)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
32177_c3_133	504	2424	64	195		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3316436_f1_19	505	2425	462	1389	331	7.4e-30
Protein name					s Name SC ECOLI	Acc# P25535
Description				Sp. 11		
VISC PROTEIN,	·					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33632828_f3_62	506	2426	249	750	559	5.1e-54
Protein name				_	s Name	Acc#
ribose-5-phosphate	isomera	ase		gp:AF	037440	AF037440
Description						
Edwardsiella ictal cds; ribose-5-phosp					_	· · · · · ·
(iciA), putative 26 fructose 1,6-bispho phosphoglycerate ki	kDa pro sphatea	otein (ygg ldolase (f	E),putati da) genes	ve 30.6 d , complet	kDa prote	
fructose 1,6-bispho	kDa pro sphatea	otein (ygg ldolase (f	E),putati da) genes	ve 30.6 d , complet	kDa prote	
fructose 1,6-bispho phosphoglycerate ki	kDa pro sphatea nase(pgl	otein (ygg ldolase (f k) gene, p	E),putati da) genes artial cd	ve 30.6 de la completa del completa de la completa de la completa del completa de la completa del completa del completa della della completa della completa della della completa della com	kDa prote te cds; a	and
fructose 1,6-bispho phosphoglycerate ki  ORF Name	kDa prosphatea	otein (ygg ldolase (f k) gene, p <u>AAID</u>	E),putati da) genes artial cd <u>NT</u> <u>Length</u>	ve 30.6 h, completes.  AA Length  1293	kDa prote te cds; a  Score  456  Name	Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
35163902_c2_109	508	2428	627	1884	960	8.9e-	103
Protein name					s Name BA_ECOLI		<u>Acc#</u> P27299
Description							
PROBABLE TRANSPORT	ATP-BIN	DING PROT	EIN MSBA				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
35350061_c2_98	509	2429	64	195			
Protein name				Locu	s Name		Acc#
Description							
NO-HIT	<del></del>			<del></del>			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
36128378_£3_67	510	2430	124	375			
Protein name				Locu	s Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
3912568_c1_92	511	2431	525	1578	1470	1.5e-1	.50
Protein name soluble pyridine nu	cleotid	e transhyo	drogenase		s Name 159108		<u>Acc#</u> AF159108
Description							
Azotobacter vinelan gene, complete cds.	dii sol	uble pyri	dine nucl	eotide t	ranshydro	genase	(sth)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
4111008_f2_33	512	2432	94	285	240	3.2e-2	0
Protein name					s Name PA PSEAE		Acc# P95459
Description							
MAJOR COLD SHOCK PR	OTEIN C	SPA					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4500892_c1_91	513	2433	291	876	599	2.9e-58
Protein name				-	s Name	Acc#
Description				[SF 1 - 2		P03822:P46 137:P76203
HYPOTHETICAL 31.2	KD PROTI	IN IN PPS	SA-AROH IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5132667_f1_12	514	2434	368	1107	124	6.4e-05
Protein name				Locus	s Name	Acc#
mannosyltransferas	se-like p	rotein		gp:YP	S251712	AJ251712
Description						,
Yersinia pseudotu gene cluster for do gene, wzx gene, wby manC gene, wbyL gen	dhD gene, yI gene,	, ddhA ger wbyJ gene	ne, ddhB g e, wzygene	gene, ddh	Cgene, pr	t gene, wbyH
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
5859762_c2_120	515	2435	115	348		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5894501_c1_82	516	2436	551	1656	564	1.5e-54
Protein name	•				s Name 53_HAEIN	<u>Acc#</u> P44029
Description					· · · · · · · · · · · · · · · · · · ·	
HYPOTHETICAL PROT	EIN HIO6	53				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5993768_c2_122	517	2437	514	1542	305	4.6e-24
Protein name					Name	Acc# P44846
Description				[		
ORGANIC SOLVENT TO	OLERANCE	PROTEIN E	OMOLOG PR	ECURSOR		

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
7150051_c3_131	518	2438	70	213	131	1.2e-08
Protein name hypothetical protei	n APE2Î	.43		Locus pir:B	Name 72521	<u>Acc#</u> B72521
Description					<u>_</u>	
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
977055_c3_129	519	2439	327	984	505	2.7e-48
Protein name	•				Name JE_ECOLI	Acc# P75826
Description						
HYPOTHETICAL 34.4 K	D PROTE	IN IN POX	B-AQPZ IN	TERGENIC	REGION	· · · · · · · · · · · · · · · · · · ·
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
9882793_c3_128	520	2440	65	198	109	2.5e-06
Protein name				_		- "
FIOCEIII IIaile				Locus	Name	Acc#
hypothetical protei	n APE06	66		Locus pir:F		Acc# F72654
	n APE06	66		_		
hypothetical protei	n APE06	AAID	NT Length	_		
hypothetical protei  Description				pir:F	72654	F72654
hypothetical protei  Description  ORF Name  23615951_c1_12  Protein name	NTID 521	AAID 2441	Length 125	AA Length	72654 Score	F72654  Probability
hypothetical protei  Description  ORF Name  23615951_c1_12	NTID 521	AAID 2441	Length 125	AA Length	Score [182]	F72654  Probability  4.5e-14  Acc#  JC5327:PC4
hypothetical protei  Description  ORF Name  23615951_c1_12  Protein name  adhesin complex 25K	NTID 521	AAID 2441	Length 125	AA Length 378	Score [182]	F72654  Probability  4.5e-14  Acc#
hypothetical protei  Description  ORF Name  23615951_c1_12  Protein name  adhesin complex 25K protein  Description  ORF Name	NTID 521 protei	AAID 2441	Length 125 or:LecA  NT Length	AA Length    Dir:F    AA     Locus     Dir:Jo	Score  [182] S Name  25327	Probability  Acc#  JC5327:PC4 312  Probability
Description  ORF Name  23615951_c1_12  Protein name  adhesin complex 25K protein  Description	NTID 521 protei	AAID  2441  n precurs	Length 125 or:LecA	AA Length  378  Locus  pir:Jo	Score [182] S Name 25327	F72654  Probability  4.5e-14  Acc#  JC5327:PC4 312
hypothetical protei  Description  ORF Name  23615951_c1_12  Protein name  adhesin complex 25K protein  Description  ORF Name  24259425_c3_15  Protein name	NTID 521 protei	AAID  2441  n precurs  AAID  2442	Length 125 or:LecA  NT Length 165	AA Length  Locus  AA Length  AA Locus  AA Length  AB Length  Locus	Score  [182] S Name  C5327  Score  [194] S Name	Probability  Acc#  JC5327:PC4 312  Probability
hypothetical protei  Description  ORF Name  23615951_c1_12  Protein name  adhesin complex 25K protein  Description  ORF Name  24259425_c3_15	NTID 521 protei	AAID  2441  n precurs  AAID  2442	Length 125 or:LecA  NT Length 165	AA Length  AA Locus  AA Length  498	Score  [182] S Name  C5327  Score  [194] S Name	F72654  Probability  4.5e-14  Acc#  JC5327:PC4 312  Probability  2.4e-15

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33986343_±3_10	523	2443	699	2100	2411	2.9e-250
Protein name					s Name	Acc#
oligopeptidepermeas	e	31 VIIII V		gp:SP	OPPDACA	X89237
Description						
S.pyogenes DNA for	oppA,	oppB, oppC	, oppD, c	ppF, and	dacA ger	nes.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4727338_f3_9	524	2444	325	978	1264	1.0e-128
Protein name				Locus	s Name	Acc#
oligopeptidepermeas	e		<u>.</u>	gp:SP	OPPDACA	X89237
Description						
S.pyogenes DNA for	oppA,	oppB, oppC	, oppD, c	ppF, and	dacA ger	ies.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4788508_f3_11	525	2445	63	192		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6053212_f3_8	526	2446	340	1023	1398	6.3e-143
Protein name			,	Locus	s Name	Acc#
oligopeptidepermeas	e			gp:SP	OPPDACA	X89237
Description	*					
S.pyogenes DNA for	oppA,	oppB, oppC	, oppD, c	ppF, and	dacA ger	les.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12265658_c2_101	527	2447	221	666	753	1.4e-74
Protein name					s Name	Acc#
Description				sp:DP	3X_HAEIN	P43746
	~~~	- CIANAPA /				
DNA POLYMERASE III	SORONI,	r gamma/Ta	.U,			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12501562_c2_109	528	2448	214	645	254	3.5e-21
Protein name hemolysin-related	protein			Locu pir:F	s <u>Name</u> 72326	Acc# F72326
Description			<del></del>			
ORF Name	NTID	<u>AAID</u> .	NT Length 967	AA Length 2904	Score	Probability
	] [223					
Protein name					s Name TD ECOLI	Acc#
Description				ор.ги	<u> </u>	P23931:P32 982:P77350
(MUREIN HYDROLASE	D) (REGU	JLATORY PRO	OTEIN DNI	R)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12891082_f3_51	530	2450	237	714	234	1.4e-19
Protein name					s Name	Acc#
Description				Sp. 16.	HD_ECOHI	P52696:P75 761
HYPOTHETICAL TRANS	CRIPTION	AL REGULA	FOR IN MC	DC-BIOA	INTERGENI	C REGION
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13876010_±1_11	531	2451	135	408	156	7.0e-11
Protein name					s Name CR CHRVI	Acc# P25544
Description				.   55 112		
RUBISCO OPERON TRA	NSCRIPTI	ONAL REGU	LATOR			
			NT	AA		
ORF Name	NTID	AAID	<u>Length</u>	<u>Length</u>	Score	Probability
15870706_c1_68	532	2452	344	1035	1009	1.1e-101
Protein name				Locus	s Name	Acc#
Description				sp:LE	U2_ECOLI	P30127:P78
(ISOPROPYLMALATE I	SOMERASE	(ALPHA-	IPM ISOME	RASE) (II	PMI)	

ORF Name	NTID AAID NT Length	AA Score Probabilit	ΞY
175062_c1_79	533 2453 219	660 740 3.4e-73	
Protein name		Locus Name Acc	
Description			
4-HYDROXYPHENYLPYR	UVATE DIOXYGENASE, (4HPP)	D) (HPD)	
ORF Name	NTID AAID NT Length	AA Score Probabilit	·Y
19769052_c1_74	534 2454 131	396 294 2.0e-25	
Protein name		Locus Name Acc	
Description			
LYSYL-TRNA SYNTHET	ASE, (LYSINETRNA LIGAS)	E) (LYSRS)	
ORF Name	NTID AAID NT Length	AA Score Probabilit	Υ `
20178438_c1_80	535 2455 173	522 629 1.9e-61	
Protein name		Locus Name Acc	
Description			
4-HYDROXYPHENYLPYR	UVATE DIOXYGENASE, (4HPP)	D) (HPD)	
ORF Name	NTID AAID NT Length	AA Score Probabilit	<u> </u>
21729513_c3_129	536 2456 61	186	
Protein name		Locus Name Acc	<u>:#</u>
Description			
NO-HIT			
ORF Name	NTID AAID NT Length	AA Score Probabilit	Y
21738306_f2_30	537 2457 100	303 185 4.9e-14	
Protein name		Locus Name Acc sp:SECF_HAEIN P445	
Description		<u></u>	
PROTEIN-EXPORT MEM	BRANE PROTEIN SECF		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
22443750_c3_128	538	2458	201	606 166	8.1e-12
Protein name				Locus Name sp:YC54_SYNY3	Acc# P74078
Description					
HYPOTHETICAL 38.3	D PROTI	IN SLL125	4		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23572128_c1_92	539	2459	103	312 179	6.0e-13
Protein name				Locus Name sp:RADA_PSEAE	Acc# P96963
Description					
DNA REPAIR PROTEIN	RADA H	OMOLOG (DN	A REPAIR	PROTEIN SMS HOMO	LOG)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23614376_c3_119	540	2460	312	939 742	2.1e-73
Protein name				Locus Name sp:EX3_HAEIN	Acc# P44318
Description					
EXODEOXYRIBONUCLEA	SE III,	(EXONUCLE	ASE III)	(EXO III)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23994182_f1_17	541	2461	171	516 175	2.5e-13
Protein name				Locus Name	Acc#
orf1				gp:PAU39558	U39558
Description					
Pseudomonas aerugi (tolB) genes, compl			tolQ), To	olR (tolR), TolA	(tolA), and TolB
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24276625_c3_122	542	2462	293	882 337	1.7e-30
Protein name				Locus Name  Sp:YGIP ECOLI	Acc# P45463
Description					
HYPOTHETICAL TRANS	CRIPTIO	NAL REGULA	TOR IN BA	ACA-TTDA INTERGEN	IC REGION

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24406575_c1_69	543	2463	227	684	780	1.9e-77
Protein name					s <u>Name</u> UD_AZOVI	Acc# P96196
Description				4		
(ISOPROPYLMALATE IS	OMERASE	(ALPHA-	IPM ISOME	RASE)		
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
24415911_c3_115	544	2464	97	294	97	4.6e-05
Protein name				Locus	s Name	Acc#
outer membrane prot	ein H.8	precurso	c	pir:S	04157	S04157
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24417077_c3_121	545	2465	555	1668	229	9.9e-16
Protein name					Name 3X_HAEIN	Acc# P43746
Description		•				
DNA POLYMERASE III	SUBUNIT	GAMMA/TA	J,			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25584501_c2_110	546	2466	230	693		
Protein name			•	Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30198405_c2_100	547	2467	417	1254	1913	1.7e-197
Protein name					Name C ACICA	<u>Acc#</u> Q43990
Description					P	
LYSYL-TRNA SYNTHETA	SE. (LY	SINETRN	A LIGASE)	(LYSRS)		

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34406268_c1_70	548 2468	169	510		
Protein name			Locus	Name	Acc#
Description					
NO-HIT					
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3912568_c2_105	549 2469	467	1404	819	1.4e-81
Protein name				Name AC_BACFI	Acc# P27611
Description			<b></b>		
NA(+)/H(+) ANTIPOR	TER				
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3953377_c3_117	550 2470	218	657	534	3.1e-51
Protein name				Name U2_CANMA	Acc# Q00464
Description			<u> </u>		
ISOMERASE) (ALPHA-	IPM ISOMERASE) (	(IPMI)			
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3988813_£1_4	551 2471	628	1887	1391	3.5e-142
Protein name  general protein sec	cretion pathway	subunit	_	Name 179925	Acc# AF179925
SecD Description					
Citrobacter freund complete cds.	ii general prote	ein secreti	on pathwa	ay subuni	t SecDgene,
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4314068_c3_118	552 2472	359	1080	1356	1.8e-138
Protein name				Name J3 NEILA	Acc# P50180
Description					
(IMDH) (3-IPM-DH)	(FRAGMENT)				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4335328_c2_98	553	2473	62	189	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4487638_f1_1	554	2474	613	1842 1886	1.2e-194
Protein name				Locus Name sp:PPCK_CHLLI	Acc# Q08262
Description					
(PHOSPHOENOLPYRUVA	TE CARBO	OXYLASE)	(PEPCK)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
4771925_c2_94	555	2475	207	624 331	7.4e-30
Protein name				Locus Name sp:RUVA_PSEAE	Acc# Q51425
Description					
HOLLIDAY JUNCTION	DNA HEL	ICASE RUV	/A		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4866427_f3_61	556	2476	289	870 247	5.9e-21
Protein name hypothetical prote	in			Locus Name	<u>Acc#</u> \$75235
Description					·
ORF Name	NTID	AAID	NT Length	AA Score	Probability
4881338_f1_5	557	2477	287	864 420	2.7e-39
Protein name				Locus Name sp:SECF_HAEIN	Acc# P44590
Description					
PROTEIN-EXPORT MEM	BRANE PI	ROTEIN SE	ECF		<u> </u>

ORF Name NTID AAID NT AA Score Probability Length Length	
5084463_f2_28   558   2478   114   345   240   3.2e-20	
Protein name Locus Name Acc# sp:YAJC_ECOLI P1967	7
Description	
HYPOTHETICAL 11.9 KD PROTEIN IN TGT-SECD INTERGENIC REGION (ORF12)	
ORF Name NTID AAID NT AA Score Probability	
5111588_c3_116	
Protein name Locus Name Acc# fructose-1,6-bisphosphate aldolase gp:PST011927 AJ011	<del>9</del> 27
Description	
Pseudomonas stutzeri fda gene and gene encoding hypotheticalprotein.	
ORF Name NTID AAID NT AA Score Probability	
978400_c1_83	
Protein name Locus Name Acc# penicillin-binding protein 4 gp:AF156692 AF156	592
Description	
Neisseria gonorrhoeae penicillin-binding protein 4 (pbp4) gene, complete co	īs.
ORF Name NTID AAID NT AA Score Probability	
1053753_±3_63	
<u>Protein name</u> <u>Locus Name</u> <u>Acc#</u>	
putative membrane protein gp:AF150928 AF1509	28
<u>Description</u>	
Acinetobacter sp. ADP1 BenP (benP) and AreR (areR) genes, completeds; are operon, complete sequence; SalD (salD), and SalE (salE)genes, complete cds, SalR (salR), SalA (salA), putative membraneprotein, putative 2-component regulatory protein, putativehistidine kinase of 2-component regulatory system, and carbonicanhydrase homolog genes, complete cds; and dihydropyrimidinasehomolog gene, partial cds.	
ORF Name NTID AAID NT AA Score Probability	•
<u>=====================================</u>	
1058425_c2_108	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
1203450_f1_5	563	2483	530	1593 1297	3.2e-132
Protein name				Locus Name	Acc# 086233
Description					
HYPOTHETICAL PROT	EIN HIII:	26.1			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
12271926_c1_71	564	2484	143	197	1.2e-15
Protein name				Locus Name sp:YFFB_HAEIN	Acc# P44515
Description					
HYPOTHETICAL PROT	EIN HIOI	03			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
15635930_f3_61	565	2485	373	1122 842	5.2e-84
Protein name				Locus Name sp:QUEA_ECOLI	Acc# P21516
Description					
(QUEUOSINE BIOSYN	THESIS P	ROTEIN QU	EA)	<u></u>	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
16134657_f2_24	566	2486	367	1104 895	1.3e-89
Protein name				Locus Name sp:GCST_ECOLI	Acc# P27248
Description					
PROTEIN)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
197211_c1_91	567	2487	240	723 304	5.4e-27
Protein name				Locus Name	Acc#
hypothetical prot	ein			gp:ACRBDOXN	Z46863
Description					
Acinetobacter sp. ORF2 and ORF3 gene	_	obQ, sodM	, lysS, ru	bA, rubB, estB,	oxyR,ppk, mtgA,

ORF Name	NTID	AAID	<u>NT</u> Length	AA Scor	e Probability
20890660_£3_56	568	2488	121	366	
Protein name				Locus Nam	e Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Scor	e <u>Probability</u>
20915682_f1_1	569	2489	142	429 162	6.0e-12
Protein name				Locus Nam	
Description					
HYPOTHETICAL 15.6	KD PROT	EIN IN SEC	B-TDH INT	ERGENIC REGIO	N
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	e <u>Probability</u>
21759656_±1_8	570	2490	591	1776 630	1.6e-103
Protein name			.,,	Locus Nam	<del></del>
Na(+):solute sympor	rter (S	sr ramily)		pir:E70480	E70480
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Scor	e <u>Probability</u>
22353380_t1_7	571	2491	89	270	
Protein name				Locus Nam	<u>e</u> <u>Acc#</u>
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Scor	e Probability
23438426_f2_26	572	2492	961	2886 281	3.1e-299
Protein name				Locus Nam	
Description				<u> </u>	
DECARBOXYLASE) (GL	YCINE C	LEAVAGE SY	STEM P-PR	OTEIN)	

ORF Name	NTID	AAID	NT Length	AA Length	Score	Proba	ability
23444531_f2_25	573	2493	138	417	411	2.5e-	-38
Protein name				Locu	s Name		Acc#
glycine cleavage sy H:aminomethyl carrie decarboxylase comple	r prote	in:glycine	=	pir:A	56623		A56623:S36 833:B56689 :I41231:H6
ORF Name 24391340_f3_47	<u>NTID</u>	AAID 2494	NT Length	AA Length 249	Score	Proba	ability
Protein name				Locus	s Name		Acc#
Description							
NO-HIT .							
ORF Name	NTID	AAID	NT Length	AA Length	Score		ability
24397200_£2_32	575	2495	410	1233	1330	1.0e-	135
Protein name					S Name T_HAEIN		<u>Acc#</u> P44594
Description							
TRANSGLYCOSYLASE) (	GUANINE	INSERTION	N ENZYME)		<del></del>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
25400263_c3_117	576	2496	399	1200	725	1.3e-	71
<u>Protein name</u>					Name AB_PSEFR		<u>Acc#</u> P72190
<u>Description</u>				<u> </u>			
HYPOTHETICAL 30.2 K	D PROTE	IN IN CAPE	3 3'REGIO	N			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
25562762_f2_18	577	2497	109	330	199	7.2e-	16
Protein name				Locus	Name		Acc#
glutaredoxin 3 (grx	C1) RP2	04		pir:F	71731		F71731
Description				-			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25626452_c3_125	578	2498	106	321	129	1.9e-08
Protein name					s <u>Name</u> GL_ECOLI	Acc# P76003
<u>Description</u>						
HYPOTHETICAL 12.4	KD PROT	EIN IN MI	NC-SHEA IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
282550_c3_110	579	2499	316	951	127	2.3e-05
Protein name				Locus	s Name	Acc#
hypothetical prote	in			gp:SF	R236923	AJ236923
Description						
Shewanella frigidi	marina	ifcA gene	and ORF2	(partial	and ORE	1.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29314057_c1_90	580	2500	289	870	705	1.7e-69
Protein name				Locus	s Name	Acc#
Protein name probable ion trans	porter			Locus pir:E		Acc# E75470
	porter			_		
probable ion trans	porter NTID	AAID	NT Length	_		
probable ion trans		<u>AAID</u>		pir:E	75470	E75470
probable ion trans  Description  ORF Name	NTID		Length	AA Length 432	75470 Score	E75470
probable ion trans  Description  ORF Name  29333458_f2_39	NTID		Length	Pir:E  AA  Length  432  Locus	75470 Score	E75470  Probability  2.7e-14
probable ion trans  Description  ORF Name  29333458_f2_39	NTID		Length	Pir:E  AA  Length  432  Locus	Score [196] S Name	E75470  Probability  2.7e-14  Acc#
probable ion trans  Description  ORF Name  29333458_f2_39  Protein name	NTID   [581		Length  143	AA Length 432 Locus Sp:SY	Score  [196] S Name L SYNY3	E75470  Probability  2.7e-14  Acc#
probable ion trans  Description  ORF Name  29333458_f2_39  Protein name  Description	NTID   [581	2501	Length  143	AA Length 432 Locus Sp:SY	Score  [196] S Name L SYNY3	E75470  Probability  2.7e-14  Acc#
probable ion trans  Description  ORF Name  29333458_f2_39  Protein name  Description  LEUCYL-TRNA SYNTHE	NTID  [581  TASE, (	2501	Length  143  FRNA LIGAS  NT	AA Length 432 Locus sp:SY	Score  [196] S Name L SYNY3	E75470  Probability  2.7e-14  Acc# P73274
probable ion trans  Description  ORF Name  29333458_f2_39  Protein name  Description  LEUCYL-TRNA SYNTHE  ORF Name	NTID  581  TASE, (1)	2501  LEUCINE1	Length  143  TRNA LIGAS  NT Length	AA Length 432 Locus Sp:SY  AA Length  Locus  Locus	Score  196 S Name L SYNY3 S) Score  128 S Name	Probability  2.7e-14  Acc# P73274  Probability  2.3e-05  Acc#
probable ion trans  Description  ORF Name  29333458_f2_39  Protein name  Description  LEUCYL-TRNA SYNTHE  ORF Name  30100432_f3_66	NTID  581  TASE, (1)	2501  LEUCINE1	Length  143  TRNA LIGAS  NT Length	AA Length 432 Locus Sp:SY  AA Length  Locus  Locus	Score	Probability  2.7e-14  Acc# P73274  Probability  2.3e-05

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3298257_£1_16	583	2503	178	537	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
35637_£2_30	584	2504	165	498 146	3.0e-10
Protein name unknown				Locus Name gp:AF064527	Acc# AF064527
Description		**		<del>-</del>	
Rhodocista centenar	ia PPH	(pph) ge	ne, comple	ete cds; and unkn	owngenes.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3907781_c3_126	585	2505	170	513	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3925443_c2_107	586	2506	136	411 369	6.9e-34
Protein name				Locus Name sp:RS6_ECOLI	Acc# P02358
Description					
30S RIBOSOMAL PROTE	IN S6				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4003558_f1_2	587	2507	157	474 508	1.3e-48
Protein name				Locus Name sp:DUT_ECOLI	Acc# P06968
Description					
(DUTPASE) (DUTP PYR	OPHOSPI	HATASE)			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4328443_f3_43	588	2508	151	456	393	2.0e-36
Protein name				Locus sp:SEC	Name B_ECOLI	Acc# P15040
Description						
PROTEIN-EXPORT PRO	TEIN SE	СВ				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4860762_£3_64	589	2509	268	807		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4860943_c1_89	590	2510	185	558	199	7.2e-16
Protein name				Locus		Acc#
NADPH:quinone oxid	oreduct	ase ————		gp:AF1	45234	AF145234
Description						
Arabidopsis thalia	na NADP	H:quinone	oxidoredu	ctase (NQ	R) mRNA,	complete cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4897050_t3_44	591	2511	153	462	355	2.1e-32
Protein name				Locus		Acc#
acetylglutamate ki	nase			pir:D7	0477	D70477
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
50160_c1_75	592	2512	127	384	460	1.9e-42
Protein name				Locus	Name	Acc#
haemoglobin-haptog	lobin b	inding pro	otein HhuA	gp:HIU	43198	U43198
Description						
Haemophilus influe gene, complete cds		emoglobin	-haptoglob	oin bindin	g protei	n HhuA (hhuA)

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6650718_t2_40	593	2513	82	249		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT	**			···		· · · · ·
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6822152_c3_128	594	2514	160	483	438	3.4e-41
Protein name					Name 9_ECOLI	Acc# P02418
Description						
50S RIBOSOMAL PROT	EIN L9					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
783426_£1_15	595	2515	766	2301	2022	4.8e-209
Protein name				Locus	Name	Acc#
<u>Description</u>				sp:SY	r_ecori	P07813:P78 292:P77110
LEUCYL-TRNA SYNTHE	TASE, (I	EUCINET	RNA LIGAS	E) (LEURS	5)	
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
860300_£2_29	596	2516	85	258		
Protein name				Locus	Name	Acc#
Description						
NO-HIT				v.		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
9803127_c1_88	597	2517	321	966	53	0.041
Protein name hypothetical prote	in (bpi	3' region)		Locus	Name 37397	Acc# C37397
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
9898513_c2_101	598	2518	527	1584	600	1.3e-	88
Protein name					s Name 67_HAEIN		Acc#
<u>Description</u>							Q57408:P96 344
PROBABLE TONB-DEPE	ENDENT RE	CEPTOR HI	1567 PREC	URSOR			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score		bility
1040887_c1_72	599	2519	301	906	601	1.8e-	58
Protein name					Name 025342		<u>Acc#</u> AB025342
Description							
Moritella marina g synthesis gene clus		omplete cd	s, simila	r to eic	osapentae	noicac	rid
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
10648402_f3_46	600	2520	395	1188	953	9.0e-	96
Protein name					s Name OF_ECOLI		<u>Acc#</u> P00888
Description				<u> </u>			
SYNTHETASE) (3-DEC	DXY-D-ARA	BINO-HEPT	ULOSONATE	7-PHOSPI	HATE SYNT	'HASE)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
10723543_f3_48	601	2521	73	222			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT		, <u>,</u>					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
10969087_£3_60	602	2522	1242	3729	2416	3.1e-	286
Protein name  DNA polymerase III	-				S Name 062919		Acc# AF062919
Description			<u>.</u>	<u>ا ا</u>			
Pseudomonas fluore	scens DN	A polymer	ase III (	dnaE) ger	ne, compl	.etecds	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
19552252_c3_122	603	2523	111	336	238	5.3e-	20
Protein name				Locus	S Name		Acc#
<u>Description</u>							D90863:AB0 01340
E.coli genomic DNA,	Kohara	clone #4	07 (52.4-5	2.8 min.	<b>7.</b>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
20180387_c2_104	604	2524	208	627	117	1.0e-	05
Protein name					S Name 66_HAEIN		<u>Acc#</u> P43988
Description							
HYPOTHETICAL PROTEI	N HI036	6 PRECURS	OR			~~	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
20355003_f2_25	605	2525	173	522			
Protein name				Locus	s Name		Acc#
Description							
по-ніт							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
20486501_c3_115	606	2526	154	465	239	4.1e-	20
Protein name hypothetical protei	n PH033	6		Locus pir:E	Name 71140		<u>Acc#</u> E71140
Description				J [			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
2120263_c3_114	607	2527	200	603	217	8.9e-	18
Protein name				•	Name		Acc# P11666
Description							
(F286)		·					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22131925_c3_124	608	2528	387	1164	1176	2.1e-119
Protein name			<del>-</del>	_	S Name U67933	Acc# U67933
Description						
Providencia stuart	ii AarC	(aarC) ge	ne, compl	ete cds.		
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
24219200_f3_59	609	2529	413	1242	872	3.5e-87
Protein name					s Name FD_HAEIN	Acc# P44683
Description				<u></u>		
HYPOTHETICAL PROTE	IN HIO3	96				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
24412678_f2_27	610	2530	253	762	291	1.3e-25
Protein name					Name H2_VIBCH	Acc# P52021
Description						
RIBONUCLEASE HII,	(RNASE I	HII) (FRAGI	MENT)			
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
24423250_f3_53	611	2531	67	204		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25573802_f1_4	612	2532	446	1341	456	4.2e-43
Protein name	de syntl	ase,		Locus	Name 54180	Acc# E64180
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29407800_c1_80	613	2533	403	1212	1125	5.4e-114
Protein name					s Name	Acc#
<u>Description</u>						Q51385:Q51 525
HYPOTHETICAL 41.7	KD PROTI	EIN IN PIL	F-NDK INT	ERGENIC	REGION (C	ORF1)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3145438_c1_69	614	2534	485	1458	644	1.3e-66
Protein name				Locus	s Name	Acc#
unknown				gp:AF	003741	AF003741
Description						
Escherichia coli	CFT073 pa	athogenici	ty island	gene, co	omplete c	ds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33882816_c1_79	615	2535	273	822	243	1.2e-41
Protein name					s Name	Acc#
Description				sp:IF	CB_ECOLI	P39199:P78 252:P76939
(EC 2.1.1.72)	<u> </u>					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3906293_f1_3	616	2536	110	333	146	3.0e-10
Protein name					s Name AL ECOLI	Acc# P76053
Description						
HYPOTHETICAL 21.5	KD PROT	EIN IN OGT	'-DBPA IN'I	ERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3907568_c2_105	617	2537	395	1188	386	1.1e-35
Protein name					s Name GL ECOLI	Acc# P77774
Description				DP. 11		
HYPOTHETICAL 41.9	KD PROT	EIN IN XSE	A-HISS IN	TERGENIC	REGION	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
3912818_c3_125	618	2538	425	1278 1077	6.6e-109
Protein name				Locus Name sp:SYH_ECOLI	Acc# P04804
Description					
(HISRS)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3945293_c3_113	619	2539	328	987 696	1.5e-68
Protein name				Locus Name sp:SOHB_HAEIN	Acc# P45315
Description					
POSSIBLE PROTEASE	SOHB,				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3946892_c3_126	620	2540	280	843 164	6.3e-12
Protein_name				Locus Name sp:Y370_HAEIN	Acc# P43989
Protein name  Description					
	IN HIO3	70			
Description	IN HIO3	70 AAID	<u>NT</u> Length		
Description HYPOTHETICAL PROTE				sp:Y370_HAEIN	P43989
Description  HYPOTHETICAL PROTE	NTID	AAID	Length	sp:Y370_HAEIN  AA Length Score 618 508  Locus Name	Probability  1.3e-48  Acc#
Description  HYPOTHETICAL PROTE  ORF Name  3946917_t3_64	NTID	AAID	Length	sp:Y370_HAEIN  AA Length 518 508	Probability  1.3e-48  Acc#
Description  HYPOTHETICAL PROTE  ORF Name  3946917_t3_64	NTID	AAID	Length	sp:Y370_HAEIN  AA Length Score 618 508  Locus Name	Probability  1.3e-48  Acc#
Description  HYPOTHETICAL PROTE  ORF Name  3946917_t3_64  Protein name	NTID	AAID	Length	sp:Y370_HAEIN  AA Length Score 618 508  Locus Name	Probability  1.3e-48  Acc#
Description  HYPOTHETICAL PROTE  ORF Name  3946917_t3_64  Protein name  Description	NTID	AAID	Length	sp:Y370_HAEIN  AA Length Score 618 508  Locus Name	Probability  1.3e-48  Acc# P44321  Probability
Description  HYPOTHETICAL PROTE  ORF Name  3946917_f3_64  Protein name  Description  GLYCOSIDASE) (TAG)	NTID 621	<u>AAID</u> [2541	Length 205	sp:Y370_HAEIN  AA Score Length 508  Locus Name sp:3MGA_HAEIN	Probability  1.3e-48  Acc# P44321
Description  HYPOTHETICAL PROTE  ORF Name  3946917_f3_64  Protein name  Description  GLYCOSIDASE) (TAG)  ORF Name	NTID 621 NTID 622	AAID  AAID  AAID  2542	Length  205  NT Length	Sp:Y370_HAEIN  AA Score  618 508  Locus Name  Sp:3MGA_HAEIN  AA Score	Probability  1.3e-48  Acc# P44321  Probability

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4181537_c1_88	623 2543	473	1422	952	2.2e-144
Protein name			Locus	s Name	Acc#
			sp:YF	GK_ECOLI	P77254
Description					<del></del>
HYPOTHETICAL GTP-E	BINDING PROTEIN IN	XSEA-HIS	S INTERG	ENIC REGI	ON
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4460938_t2_37	624 2544	271	816	664	3.8e-65
Protein name			Locus	s Name	Acc#
O-acetylserine syn	thase		gp:AF	010139	AF010139
Description					
Azotobacter vinela cysE2, iscS, iscU, partial cds.					
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5114700_c1_86	625 2545	77	234		
Protein name			Locus	Name	Acc#
Description					
NO-HIT					
ORF Name	NTID AAID	<u>NT</u> Length	AA Length	Score	Probability
52138_c1_91	626 2546	109	330	199	1.2e-15
Protein name			Locus	Name	Acc#
solanesyl diphosph	ate synthase		gp:AB	001997	AB001997
Description	<u>-</u>				
Rhodobacter capsul	atus DNA for sola	nesyl dip	hosphate	synthase	c,complete cds.
ORF Name	NTID AAID	<u>NT</u> Length	AA Length	Score	Probability
6140680_f2_36	627 2547	300	903	187	1.9e-29
Protein name		*	Locus	Name	Acc#
hypothetical prote	in b2532		pir:C	55030	C65030
Description			<del>-</del> L		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
648425_c1_90	628	2548	87	264		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
10378_c2_184	629	2549	85	258	251	2.2e-21
Protein name				_	s Name	Acc#
cold shock protein,	CSPA			gp:VC	CSPA	Y11908
Description						
V.cholerae cspA ger	ie.					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1063510_c2_198	630	2550	175	528	208	1.5e-15
Protein name				Locus	Name	Acc#
uridylyl transferas	е	444,1403		gp:AB	024601	AB024601
Description						
Pseudomonas aerugin tetrahydrodipicolina	_	_		complete	e cds, st	crain PAO1.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1175012_c1_174	631	2551	411	1236	929	3.2e-93
Protein name	•			Locus	Name	Acc#
acetate kinase				pir:B'	75254	B75254
Description				·		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
11988812_c1_171	632	2552	161	486		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12532562_c3_232	633	2553	299	900		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT					······································	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12542005_f3_130	634	2554	92	279		
Protein name				Locus	s Name	Acc#
Description			•			
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13001052_f1_50	635	2555	62	189		
<u>Protein name</u>				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13852337_c2_202	636	2556	351	1056	515	2.3e-49
Protein name					s Name	Acc#
				sp:AP	BE_HAEIN	P44550
Description				-		
THIAMINE BIOSYNTHE	SIS LIP	OPROTEIN A	PBE PRECU	RSOR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
14225300_c3_224	637	2557	123	372	440	2.1e-41
<u>Protein name</u>					Name	Acc#
PII-protein				gp:AV	J91902	U91902
Description						
Azotobacter vinelar protein (amtB) gene			(glnB) an	d methyla	ammoniumt	ransport

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
15628127_£3_98	638	2558	72	219	
Protein name				Locus Name	Acc#
Description					
NO-HIT			<del></del>		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
15630192_c3_217	639	2559	162	489 270	2.0e-25
Protein name				Locus Name	Acc#
Description				sp:UP04_ECOL1	P39169:P76 624:P77022 :P77023
UNKNOWN PROTEIN FR	OM 2D-P	AGE (SPOT	LM6)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
16597790_£3_131	640	2560	353	1062 232	2.6e-19
Protein name				Locus Name sp:NUC1_CUNE	Acc# P81203
Description					
NUCLEASE C1,					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16603411_f2_85	641	2561	319	960 597	4.8e-58
Protein name				Locus Name	<u>Acc#</u>
				sp:YF56_HAEIN	P45250
Description					
PUTATIVE 2-HYDROXY	ACID DE	HYDROGENAS			
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length Score	Probability
17036428_£3_138	642	2562	62	189	
Protein name				Locus Name	Acc#
Description					
NO-HIT					

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
19564458_£2_55	643	2563	212	639	192	4.0e-15
Protein name					s Name	Acc#
probable glpG prote	in			pir:D	71258	D71258
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
21766930_f3_100	644	2564	610	1833	554	1.7e-53
Protein name				Locu	s Name	Acc#
hypothetical protei	n.			pir:S	75944	S75944
Description			<del></del>		<del></del>	
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
22035932_c2_209	645	2565	416	1251	515	2.3e-49
Protein name				Locu	s Name	Acc#
B1306.06c protein				gp:ML	B1306	Y13803
Description						
Description						
	e cosm	id B1306 I	ONA.			
Mycobacterium lepra	e cosm	ld B1306 I		ΔΔ		
	NTID	AAID	NA.  NT  Length	AA Length	Score	Probability
Mycobacterium lepra			NT		<u>Score</u>	Probability 2.0e-185
Mycobacterium lepra	NTID	AAID	NT Length	Length 1632	1799	2.0e-185
Mycobacterium lepra ORF Name 22775251_c1_144	NTID 646	<u>AAID</u>	NT Length	Length 1632	1799 s Name	
Mycobacterium lepra ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-a	NTID 646	AAID 2566 large	NT Length 543	Length 1632 Locu	1799 s Name	2.0e-185 Acc# E64729:S14
Mycobacterium lepra ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-achain	NTID 646	AAID 2566 large	NT Length 543	Length 1632 Locu	1799 s Name	2.0e-185 <u>Acc#</u> E64729:S14 385:S40590
Mycobacterium lepra ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-a	NTID 646	AAID 2566 large	NT Length 543	Length 1632 Locu	1799 s Name	2.0e-185 Acc# E64729:S14
Mycobacterium lepra ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-achain	NTID 646	AAID 2566 large	NT Length  543  large  NT Length	Length 1632 Locu	1799 s Name	2.0e-185 <u>Acc#</u> E64729:S14 385:S40590
Mycobacterium lepra  ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-achain Description	NTID 646 se, III	AAID  2566  large  thase III	NT Length 543	Length 1632 Locu pir:Y	1799 s Name CEC31	2.0e-185 Acc# E64729:S14 385:S40590 :A01113:I4
Mycobacterium lepra  ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-achain  Description  ORF Name	NTID 646 se, III cid syr	AAID  2566  large thase III  AAID	NT Length  543  large  NT Length	Length 1632 Locu pir:Y  AA Length 540	1799 s Name CEC31 Score	2.0e-185  Acc#  E64729:S14 385:S40590 :A01113:I4  Probability  1.8e-41
Mycobacterium lepra  ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-achain Description  ORF Name  22890836_c2_214	NTID 646 se, III cid syr	AAID  2566  large thase III  AAID	NT Length  543  large  NT Length	Length Locu pir:Y  AA Length  540 Locu	1799 s Name CEC31 Score 415	2.0e-185  Acc#  E64729:S14 385:S40590 :A01113:I4  Probability
Mycobacterium lepra  ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-achain Description  ORF Name  22890836_c2_214	NTID 646 se, III cid syr	AAID  2566  large thase III  AAID	NT Length  543  large  NT Length	Length Locu pir:Y  AA Length  540 Locu	1799 s Name CEC3I  Score 415 s Name	2.0e-185  Acc#  E64729:S14 385:S40590 :A01113:I4  Probability  1.8e-41  Acc#
Mycobacterium lepra ORF Name  22775251_c1_144  Protein name acetolactate synthachain:acetohydroxy-achain Description  ORF Name  22890836_c2_214  Protein name	NTID 646 se, III cid syr	AAID  2566  large thase III  AAID  2567	NT Length  543  large  NT Length  179	Length Locu pir:Y  AA Length 540 Locu gp:AH	Score 415 S Name U56832	2.0e-185  Acc#  E64729:S14 385:S40590 :A01113:I4  Probability  1.8e-41  Acc# U56832

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
23595787_c3_220	648	2568	150	453		
Protein name				Locus 1	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
23718750_f2_93	649	2569	202	609	432	1.5e-40
Protein name				Locus I		Acc# P44633
Description						
JUNCTION NUCLEASE	RUVC) (I	HOLLIDAY	JUCTION RE	SOLVASE RU	VC)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
23727181_c2_215	650	2570	330	993		
——————————————————————————————————————						
Protein name			<u> </u>	Locus 1	Name	Acc#
Protein name Description				Locus 1	<u>Name</u>	Acc#
				Locus 1	<u>Name</u>	Acc#
Description	NTID	AAID	NT Length	AA	Name core	Acc#
Description NO-HIT	<u>NTID</u>	<u>AAID</u> 2571	<del></del>	AA Length		
Description  NO-HIT  ORF Name  23939008_c1_161  Protein name	651		Length	AA Solution 1972  Locus 1972	core	Probability 1.0e-09 Acc#
Description  NO-HIT  ORF Name  23939008_c1_161  Protein name hypothetical prote	651		Length	AA Length So 372	core	Probability
Description  NO-HIT  ORF Name  23939008_c1_161  Protein name	651		Length	AA Solution 1972  Locus 1972	core	Probability 1.0e-09 Acc#
Description  NO-HIT  ORF Name  23939008_c1_161  Protein name hypothetical prote	651		Length	AA So	core	Probability 1.0e-09 Acc#
Description  NO-HIT  ORF Name  23939008_c1_161  Protein name hypothetical prote  Description	651	2571	Length  123  NT	AA So	141   Name   511	Probability  1.0e-09  Acc#  T10511
Description  NO-HIT  ORF Name  23939008_c1_161  Protein name hypothetical prote  Description  ORF Name	651 in NTID	2571 AAID	Length  123  NT Length	AA Length  Dir:T10  AA Length	Core 141 Name 511 Core	Probability  1.0e-09  Acc#  T10511
Description  NO-HIT  ORF Name  23939008_c1_161  Protein name hypothetical prote  Description  ORF Name  23942567_c2_212	651 in NTID	2571 AAID	Length  123  NT Length	AA Length  Dir:T10:  AA Length  201	Core 141 Name 511 Core	Probability  1.0e-09  Acc# T10511  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23963325_f3_123	653	2573	403	1212	879	6.3e-88
Protein name					Name DH_ECOLI	Acc# P42593
Description						
A REDUCTASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24353427_c2_200	654	2574	546	1641	2074	1.5e-214
Protein name					S Name 60_YEREN	<u>Acc#</u> P48219
Description						
60) (CROSS-REACTING	PROTE	N ANTIG	EN)	<u></u>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24612761_c3_246	655	2575	259	780	467	2.9e-44
<u>Protein name</u>					Name FC_HAEIN	Acc# P44827
Description						
HYPOTHETICAL PROTEI	N HI069	94		.,		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24648387_c1_170	656	2576	282	849	926	6.6e-93
Protein name thymidylate synthas	e		<del> </del>	Locus gp:L78	Name	<u>Acc#</u> L78665
Description				] [21		
Methylobacillus fla protein (orf-1), hom thymidylate sythase	noserine	dehydro	ogenase (ho	m), andth		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
24796885_c2_197	657	2577	268	807	801	1.2e-79
Protein name				-	Name PM ECOLI	Acc# P07906
Description						
METHIONINE AMINOPER	TIDASE,	(MAP)	(PEPTIDASE	M)		

ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25390711_c1_153	658	2578	414	1245	593	2.8e-60
Protein name					s <u>Name</u> TY_ECOLI	Acc# P17802
Description						
A/G-SPECIFIC ADEN	INE GLYCOSY	LASE,				
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25412907_£1_37	659	2579	95	288	74	0.044
<u>Protein name</u>				Locus	s Name	Acc#
hypothetical prote	ein			gp:AP	000363	AP000363
Description						
Bacteriophage VT2	-Sa, comple	te genom	ne sequen	.ce.		
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25566577_f1_23	660	2580	83	252		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID A	AID	NT	<u>AA</u>	Score	Probability
			Length	Length		
25585263_f2_64	661	2581	311	936	790	1.7e-78
Protein name diaminopimelate er	nimerage			_	Name	Acc#
				pir:S	01913	B65185:S30
<u>Description</u>						699:S01913 :A37841:S2
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26213386_c3_242	662	2582	404	1215	393	2.0e-36
Protein name				Locus	s Name	Acc#
				sp:UB	IH_ECOLI	P25534
Description				h		
UBIH PROTEIN,						

ORF Name	NTID	AAID	<u>NT</u> Length	Length Score	Probability
26214428_c2_201	663	2583	296	891 719	5.7e-71
Protein name				Locus Name sp:LGT_SALTY	Acc# Q07293
Description					<del></del>
PROLIPOPROTEIN DIA	CYLGLYC	ERYL TRAN	SFERASE,		
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
29303578_£2_74	664	2584	69	210	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
29380307_c2_199	665	2585	166	501	
Protein name				Locus Name	Acc#
Description					
NO-HIT			,		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
30268885_c3_234	666	2586	63	192	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
33626535_£1_51	667	2587	141	426	
Protein name				Locus Name	Acc#
Description					
NO-HIT		····			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34416075_c3_231	668	2588	803	2412	1247	6.3e-127
Protein name					s <u>Name</u> RX_AZOVI	Acc# P36223
Description						
TRANSFERASE) (URIDY	LYL REM	MOVING ENZ	YME)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35425918_c3_241	669	2589	176	531	302	8.7e-27
Protein name				Locus	s Name	Acc#
dihydrofolate reduc	tase,			pir:S	52336	S52336
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36051061_f2_78	670	2590	327	984	930	2.5e-93
Protein name				Locus	s Name	Acc#
probable 2				pir:G	70875	G70875
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
3910943_c1_146	671	2591	343	1032	1278	3.3e-130
Protein name				_	Name	Acc#
ketol-acid reductoi	someras	e		gp:AF	125563	AF125563
Description						
Neisseria meningiti carbomyltransferase complete cds.						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3914002_f3_114	672	2592	253	762		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3939063_f2_89	673	2593	469	1410	627	3.2e-61
Protein name					s Name RD_ECOLI	Acc# P14900
Description						
ADDING ENZYME)				-		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3947713_f1_5	674	2594	260	783	594	1.0e-57
Protein name					Name AA_HAEIN	Acc# P43908
Description			•	<del></del>		<del></del>
HYPOTHETICAL PROT	EIN HI09	34				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3954638_f2_63	675	2595	438	1317	1067	7.5e-108
Protein name				-	Name	Acc# P19572
Description				<b>P</b> 1 3 5		
DIAMINOPIMELATE D	ECARBOXY	LASE, (DAP	DECARBOX	YLASE)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3962885_c3_219	676	2596	181	546	486	2.8e-46
Protein name				Locus	Name	Acc#
acetolactate synt chain:acetohydroxy	•		gmall	pir:Y	СЕСЗН	F64729:S14
chain	acra by.					386:S40591
Description				_		:A01114:PS
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
3992035_c1_175	677	2597	500	1503	1163	4.6e-127
Protein name					Name A_HAEIN	Acc# P45107
Description				<u> </u>		
PHOSPHATE ACETYLT	RANSFERAS	E, (PHOSP	HOTRANSAC	ETYLASE)		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4140881_f1_40	678	2598	467	1404	710	5.1e-70
Protein name NorM				Locus gp:AB0	Name 10463	Acc# AB010463
Description						
Vibrio parahaemol	yticus g	ene for N	orM, compl	ețe cds.		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4147562_c1_159	679	2599	523	1572	1209	6.7e-123
Protein name				<u>Locus</u> sp:YIF	Name B_HAEIN	Acc# P45049
Description						
HYPOTHETICAL PROT	EIN HIII	17				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4471887_f3_135	680	2600	415	1248	637	2.8e-62
Protein name				Locus		Acc#
FtsW		<del></del>		gp:AF1	.23260	AF123260
Description						
Coxiella burnetii	FtsW (i	tsW) gene	, complete	cds.		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4720313_c1_180	681	2601	149	450	161	1.2e-10
Protein name				Locus		Acc#
				sp:MET	E_ECOLI	P25665
Description	_					
(COBALAMIN-INDEPE	ENDENT ME'	THIONINE				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4770887_f2_72	682	2602	176	531	130	2.7e-14
Protein name hypothetical prot	ein		· · · · · · · · · · · · · · · · · · ·	Locus gp:SSU		Acc# Y18930
Description						
Sulfolobus solfat	aricus 2	31 kb gen	omic DNA f	ragment,	strain P	22.

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
5100010_±1_7	683 2603	333	1002 581	2.4e-56
Protein name	,		Locus Name	Acc# P44818
Description				
INTEGRASE/RECOMBIN	NASE XERC			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
5275250_c3_235	684 2604	202	609 328	1.5e-29
Protein name			Locus Name	Acc#
Trp repressor bind	ling protein	,	gp:AF067083	AF067083
Description				
Vitreoscilla sp. o repressor binding p	<del>-</del>			· -
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
593802_£2_71	685 2605	506	1521 433	1.1e-40
Protein name			Locus Name	<u>Acc#</u> P44608
Description				
RIBONUCLEASE BN,	(RNASE BN)			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
5970933_f1_15	686 2606	105	318 249	3.6e-21
Protein name			Locus Name	Acc#
unknown protein			gp:MSGTCWPA	M15467
Description				
M.tuberculosis 65	kDa antigen (cell	wall pro	tein a) gene.	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
5988327_c3_227	687 2607	927	2784 1588	4.6e-163
Protein name			Locus Name	Acc#
Description			sp:HEPA_ECOLI	P23852:P75
<del></del>	SOCIATED PROTEIN (	A THO INCIDENT	DENIP PERIOTON CE DE	633

ORF Name	NTID	AAID	<u>NT</u> Length	Length So	core l	Probability
6488910_f2_94	688	2608	66	201	85	0.0050
Protein name				Locus I		Acc# P32139
Description				<u> </u>		
HYPOTHETICAL 34.	KD PROTE	EIN IN GL	NA-RBN INT	ERGENIC RE	GION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length		Probability
6516885_c2_185	689	2609	213	642	140	3.5e-09
Protein name putative membrane	e protein.			Locus 1		Acc# AL133213
Description				<del></del>		<del></del>
Streptomyces coe	licolor co	osmid 6D7	•			
ORF Name	NTID	AAID	NT Length	Length -		Probability
665876_c3_245	690	2610	552	1659	215	1.1e-16
Protein name				Locus I		<u>Acc#</u> Q05146
Description						
OUTER MEMBRANE P	ROTEIN A I	PRECURSOR	•			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>Length</u> —		Probability
8575_c3_236	691	2611	140	423	343	1.0e-31
Protein name				Locus 1		<u>Acc#</u> 033499
Description				<del></del>		
10)						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core I	Probability
859388_f2_86	692	2612	342	1029	537	[.le-51
Protein name				Locus 1		Acc# P45204
Description				L		
(FPP SYNTHASE)		<del></del> ,				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
970625_c1_145	693	2613	68	207	197	1.1e-14
Protein name					s Name VI_ECOLI	Acc#
<u>Description</u>						P00893:P78 045
III) (ACETOHYDROX	Y-ACID S	YNTHASE II	I LARGE S	SUBUNIT)	(ALS-III)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
9957828_f2_88	694	2614	149	450	85	0.00086
Protein name				Locus	s Name	Acc#
transposase				gp:CE	rc2	
Description						X59156:S88 451
Caenorhabditis ele	gans tra	ansposon T	c2.			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21736625_c2_11	695	2615	290	870	508	1.3e-48
Protein name					S Name AH_HAEIN	Acc# P44751
Description				<u> </u>		
(DIADENOSINE TETRA	APHOSPHA:	rase)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
26053825_f1_4	696	2616	528	1587	1002	5.8e-101
Protein name					Name AB_ECOLI	Acc# P03005
Description				<u> </u>		
REPLICATIVE DNA H	ELICASE,					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33984677_±3_6	697	2617	382	1146	555	1.4e-53
Protein name biosynthetic alani	ne racen	nase		_	Name 165882	Acc# AF165882
Description				<u> </u>		
Pseudomonas aerug cds.	inosa bio	osynthetic	alanine	racemase	(alr) ge	ne, complete

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
5181430_c3_14	698 2618	327	984 617	3.6e-60
Protein name			Locus Name	Acc# P19624
Description				
PYRIDOXAL PHOSPHATE	BIOSYNTHETIC	PROTEIN PDX	(A	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
7680_c3_15	699 2619	295	888 603	1.1e-58
Protein name			Locus Name sp:KSGA_ECOL	Acc# P06992
Description				
DIMETHYLTRANSFERASE	3)			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
10547156_c1_79	700 2620	1096	3291 4176	0.0
Protein name			Locus Name	Acc#
carbamoylphosphate	synthetase larg	ge subunit	gp:PAU81259	1181250 127
Description				U81259:L27 528
Pseudomonas aerugir cds, carbamoylphosph synthetase large sub gene, partial cds.	nate synthetase	small subu	nit (carA) andca	rbamoylphosphate
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
11915953_c1_76	701 2621	138	417 312	7.6e-28
Protein name probable oxidoreduc	tase		Locus Name	<u>Acc#</u> T35853
Description				
ORF Name [13947152_f2_37	NTID AAID	NT Length	AA Length Score	Probability
Protein name	J		Locus Name	Acc#
Description				
NO-HIT		·		

ORF Name	NTID	AAID	NT Length	AA Length	Probability
14094052_f1_20	703	2623	111	336 160	9.7e-12
Protein name				Locus Name sp:YCCK_ECOLI	
Description					P45572:P75 878
HYPOTHETICAL 12.4	KD PROT	EIN IN HE	LD-SERT IN	TERGENIC REGION	<u> </u>
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16601077_f2_30	704	2624	93	282 103	1.1e-05
Protein name				Locus Name	Acc#
hypothetical prot	ein			gp:SSU18930	Y18930
Description					
Sulfolobus solfat	aricus 2	81 kb gen	omic DNA f	ragment, strain	P2.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19562686_f3_66	705	2625	309	930 530	6.0e-51
Protein name				Locus Name	Acc#
Description				sp:CBL_ECOLI	Q47083:P76
TRANSCRIPTIONAL R	EGULATOR	CBL			
			NT	AA	
ORF Name	NTID	AAID	Length	Length Score	Probability
19571925_c3_117	706	2626	70	213	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20313326_f2_40	707	2627	65	198	
Protein name				Locus Name	Acc#
Description					
NO-HIT					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabili	ty
20488827_f1_10	708	2628	558	1677	1638	2.3e-168	
Protein name sulfite reductase					s Name 026066	Ac AF0	<u>c#</u> 26066
Description							
Pseudomonas aerugi	nosa su	lfite redu	ctase (cy	sI) gene	, complet	e cds.	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabili	ty
20953402_c2_103	709	2629	328	987	213	3.2e-15	
Protein name					s <u>Name</u> PB_HAEIN	Ac P44	
Description							
OUTER MEMBRANE ANT	IGENIC I	JIPOPROTEL	N B PRECO	RSOR			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabili	ty
21907078_f3_55	710 ·	2630	66	201			•
Protein name				Locus	s Name	Ac	<u>c#</u>
Description							
NO-HIT							
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probabili	έy
22766067_f1_21	711	2631	478	1437	96	0.012	
Protein name				-	s Name DF_MYCGE	Ac	<u>c#</u>
Description						P472 330	54:Q49
POSSIBLE THIOPHENE	AND FU	RAN OXIDAT	ION PROTE	IN THDF			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	-Y
24251510_c2_88	712	2632	364	1095	746	7.8e-74	
Protein name				Locus	s Name	Ace	<u>c#</u>
hypothetical prote	in Rv362	9C		pir:F	70561	F70	561
Description							

ORF Name	NTID	AAID	Length	Length	Score	Probability
24259555_£2_39	713	2633	219	660	375	1.6e-34
Protein name					s Name	Acc#
similar to glutat	nione-s-t	ransieras	e ————	gp:AF	036940	AF036940:A
Description						F081362
Pseudomonas sp. U	2 plasmic	pwwuz,	Eerredoxin	n reducta	se	
(nagAa), salicylate						
(nagG), salicylate-						
(nagAb), naphthale naphthalene dioxyg						
dihydrodiol dehydr						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24801562_c2_82	714	2634	554	1665	506	8.0e-73
Protein name			<u> </u>	Locu	s Name	Acc#
permease for AmpC	beta-lac	tamase e	pression		082985	AF082985
Description				<u> </u>		
Pseudomonas aerug (ampG) gene, compl				a-lactam	ase expr	essionAmpG
(ampa) gene, compr	————	and annin				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25448412 c2_95	715	2635	98	297	186	1.7e-14
				<u> </u>		
Protein name				_	s Name	Acc#
unknown				] gp:Al	033858	AF033858
Description						
Pediococcus pento	saceus si	rain ATC	C43200 pla	asmid pMD	136, com	pleteplasmid
sequence.						
	NIMITO	7 7 TD	NT	AA	Caoro	Drobabilitu
ORF Name	NTID	AAID	Length	Length	Score	Probability
29429590_f2_44	716	2636	162	489	241	2.5e-20
Protein name				Locu	s Name	Acc#
				sp:VI	DLD_HELPY	
Description				<u> </u>	·	
PROTEIN VOLD						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30181587_£1_9	717	2637	76	231	43	0.037
Protein name				Locus	s Name	Acc#
bone morphogenetic	protein	. 2		pir:A	61387	A61387
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3129637_c2_94	718	2638	323	972	577	6.3e-56
Protein name				Locus	s Name	Acc#
mrr restriction sys	tem pro	tein		pir:F	75508	F75508
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31375212_f3_54	719	2639	615	1848	1278	3.3e-130
Protein name				Locus	s Name	Acc#
				sp:YF	BQ_ECOLI	P77727
Description						
PROBABLE AMINOTRANS	FERASE	YFBQ,				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
32878_f2_25	720	2640	349	1050	176	1.2e-10
Protein name		· ·			s Name	Acc#
Description				gp:EC	OTIOK	D10483:J01 597:J01683 :J01706:K0
E.coli K12 genome,	0-2.4mi	n. region	•			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33788387_c3_108	721	2641	66	201	87	0.00053
Protein name				Locus	s Name	Acc#
hypothetical protei	n SPCP3	1B10.02		pir:T	41692	T41692

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
3915688_c1_78	722 2642	200	603 482	7.4e-46
Protein name			Locus Name	Acc# P43984
Description				
HYPOTHETICAL PROT	EIN HIO318			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
40875_t1_18	723 2643	126	381 189	8.2e-15
Protein name			Locus Name sp:YHEN_ECOLI	Acc# P45532
Description				<del></del>
HYPOTHETICAL 13.6	KD PROTEIN IN R	PSL-FKPA IN	TERGENIC REGION	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
4490902_c2_101	724 2644	166	501 522	4.3e-50
Protein name			Locus Name sp:GREA_ECOLI	Acc#
Description			<u> </u>	P21346:P78
GREA)			-1-1	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
4536668_c3_107	725 2645	940	2823 1554	1.9e-159
Protein name			Locus Name	Acc#
Description			sp:GLNE_ECOLI	P30870:P78
SYNTHETASE ADENYL	YLTRANSFERASE) (	ATASE)		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4572125_c1_72	726	2646	311	936	820	1.1e-81
Protein name unknown				_	s Name U63816	Acc# U63816
Description						
Pseudomonas aerugin heptosyltransferase homolog (waaC),gluco andunknown protein (	homolog syltran	(waaF),l sferase I	ipopolysa homolog	ccharide (waaG),	heptosyl RfaP prot	transferase I cein (waaP),
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5084827_t3_48	727	2647	163	492	193	1.5e-14
Protein name				Locus	s Name	Acc#
Description				sp:SU	RA_ECOLI	P21202:P75 630
SURA), (PPIASE) (RO	TAMASE (	2)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5287555_c2_102	728	2648	125	378	125	5.0e-08
Protein name					s <u>Name</u> 53_HAEIN	<u>Acc#</u> P44139
Description						
HYPOTHETICAL PROTEI	N HI125	3				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
5322062_c1_69	729	2649	265	798	449	2.3e-42
Protein name				-	S Name EB_ECOLI	Acc# P12282
Description					•	
MOLYBDOPTERIN BIOSY	NTHESIS	MOEB PRO	rein			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
6822208_c2_83	730	2650	296	891	502	5.6e-	48
Protein name  Description				-	s Name MK_ECOLI		<u>Acc#</u> P37186:Q46
Description							754
HEMK PROTEIN							
ORF Name	NTID	AAID	NT Length	AA Length	Score		bility
6829637_f2_38	731	2651	455	1368	926	6.6e-	93
Protein name					s Name SG_ECOLI		Acc#
Description					,	<del></del>	P11098:P76 685
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
7117182_c3_114	732	2652	115	348	200	5.6e-	16
Protein name				_	s Name		Acc#
				] gp:AF	033858		AF033858
Description		1 maa			1 2 6	11	
Pediococcus pentosa sequence.	.ceus st	rain ATCC	13200 pla	smid pMD.			asmid
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
7281562_c1_77	733	2653	416	1251	739	6.4e-	130
Protein name					s Name RA_PSEAE		Acc# P38098
Description				·			
PHOSPHATE SYNTHETAS	E GLUTA	MINE CHAIL	4)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	bility
953181_c3_106	734	2654	450	1353	449	2.3e-	42
Protein name					s Name 6S_MYCTU		<u>Acc#</u> P96936
Description				L	<del>_</del>		
HYPOTHETICAL 54.8 K	D PROTE	IN CY20H1	).28C			<u>-</u> .	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
978465_±1_19	735	2655	102	309	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16985642_c2_43	736	2656	69	210 127	9.7e-08
<u>Protein name</u>				Locus Name	
Description					
GLUTAMATE RACEMASE	,		<del></del>	-	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
189186_c3_54	737	2657	233	702	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
23628411_±1_6	738	2658	295	888 379	6.1e-35
<u>Protein name</u>				Locus Name	<del></del>
Description					
HYPOTHETICAL 30.9	D PROTI	EIN IN HEM	M-PRSA IN	TERGENIC REGIO	N N
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29494003_£2_13	739	2659	672	2019 241	6.6e-17
Protein name				Locus Name	
Description				t.	
HYPOTHETICAL 64.8 1	D PROTI	IN IN HEM	M-HEMA IN	TERGENIC REGIO	N (ORF3)

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29960761_c1_30	740	2660	468	1407	731	3.0e-72
Protein name					s Name M1_PASMU	Acc# P95525
Description						
GLUTAMYL-TRNA REDUC	TASE,	(GLUTR)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31291251_f2_15	741	2661	265	798	924	1.1e-92
Protein name				Locu gp:EC	s Name OPRS	Acc# M13174
Description						
E.coli prs gene enc	oding p	phosphorib	osylpyrop	hosphate	syntheta	ase, complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34641308_f3_23	742	2662	196	591	160	9.7e-12
Protein name					s Name LB_PSEAE	Acc# P42812
Description						
OUTER MEMBRANE LIPO	PROTEI	V LOLB PRE	CURSOR			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4869025_f3_19	743	2663	554	1665	1876	1.4e-193
Protein name			•		s Name FD_ACICA	Acc# P94132
Description				<u> </u>		
DEHYDROGENASE) (ELE	CTRON-	TRANSFERRI	NG-FLAVOP	ROTEIN D	EHYDROGEN	(ASE)
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
10552153_f1_31	744	2664	74	225	168	7.1e-12
Protein name					s <u>Name</u> 028868	Acc# AB028868
Description				<del>-</del>		
Mus musculus P4(21)	n mRNA	, partial	cds.			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
10722125_f2_65	745	2665	161	486 165	2.9e-12
Protein name				Locus Name sp:SMPA_ECOLI	Acc# P23089
Description					
SMALL PROTEIN A		<u> </u>			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
117037_f3_138	746	2666	218	657 375	1.6e-34
Protein name				Locus Name sp:Y787_HAEIN	Acc# P44052
Description					
HYPOTHETICAL PROTE	EIN HIO7	87			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
11885927_f1_51	747	2667	76	231	
Protein name				Locus Name	Acc#
Description					
NO-HIT			_		
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
12297203_c3_236	748	2668	78	237 138	2.1e-09
Protein name				Locus Name	Acc#
hypothetical prote	ein APE2	061		pir:G72510	G72510
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
12506635_f2_83	749	2669	66	201	
Protein name				Locus Name	Acc#
Description					
NO-HIT		<del></del>			

ORF Name	NTID AAID	NT Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
12697037_c2_201	750 2670	111	336 99	2.8e-05
Protein name			Locus Name	Acc# X70925
Description				
P.acidilactici gene	for d-lactate de	ehydrogen	ase.	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
13078416_c3_233	751 2671	167	504 210	4.9e-17
Protein name			Locus Name	Acc#
ribosomal-protein-s N-acetyltransferase,		a F	pir:F69768	F69768
Description			J	
ORF Name	NTID AAID	NT Length	AA Score Length 539	Probability
	732 2072	13/		
Protein name ferric uptake regul	ator		Locus Name  gp:ABDNAFUR	Acc# Y14980
Description			gp. Abbitat ok	114500
Acinetobacter bauma	nnii fur gene.			
ORF Name	NTID AAID	NT Length	AA Score	Probability
13726003_c2_196	753 2673	349	1050 736	8.9e-73
Protein name			Locus Name	Acc#
iron transport prot slr1295:protein slr1	<del>-</del>		pir:S74691	S74691
Description			J	
ORF Name	NTID AAID	NT Length 81	<u>AA</u> <u>Length</u> <u>Score</u> [246] [117]	Probability
	754 2674	81	240	3.36-07
Protein name	754 2674	01	Locus Name	Acc#
Protein name hypothetical protei				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14553432_f2_67	755	2675	303	912 647	2.4e-63
Protein name				Locus Name sp:METR_SALTY	Acc# P05984
Description					
TRANSCRIPTIONAL AC	TIVATOR	PROTEIN	METR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
15820312_c2_223	756	2676	867	2604 1893	2.2e-195
Protein name				Locus Name	Acc#
UspA2				gp:AF113611	AF113611
Description				-	
Moraxella catarrha	lis str	ain V1171	UspA2 (us	pA2) gene, compl	etecds.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16016926_f3_137	757	2677	237	714 197	3.1e-15
Protein name				Locus Name	Acc#
Protein name growth factor-resp smooth muscle:SM-20	_	protein,	vascular	Locus Name	<u>Acc#</u> A53770
growth factor-resp	_	protein,	vascular		
growth factor-resp smooth muscle:SM-20 Description	NTID	AAID	NT Length	pir:A53770  AA Length Score	A53770  Probability
growth factor-resp smooth muscle:SM-20 Description			NT	pir:A53770	A53770
growth factor-resp smooth muscle:SM-20 Description	NTID	AAID	NT Length	pir:A53770  AA Length Score	A53770  Probability
growth factor-resp smooth muscle:SM-20 Description  ORF Name  16064061_c2_187	NTID	AAID	NT Length	AA Score [1194] 738	A53770  Probability  5.5e-73  Acc#
growth factor-resp smooth muscle:SM-20 Description  ORF Name  16064061_c2_187  Protein name  Description	NTID	AAID	NT Length	pir:A53770  AA Score Length 738  Locus Name	A53770  Probability  5.5e-73  Acc#
growth factor-resp smooth muscle:SM-20 Description  ORF Name  16064061_c2_187  Protein name  Description  (EC 2.4.1)	NTID 758	<u>AAID</u> 2678	NT Length	pir:A53770  AA Score Length 738  Locus Name Sp:MURG_HAEIN	A53770  Probability  5.5e-73  Acc# P45065
growth factor-resp smooth muscle:SM-20 Description  ORF Name  16064061_c2_187  Protein name  Description	NTID	AAID	NT Length	pir:A53770  AA Score  Length 738  Locus Name  Sp:MURG_HAEIN  AA Score  Length Score	A53770  Probability  5.5e-73  Acc#
growth factor-resp smooth muscle:SM-20 Description  ORF Name  16064061_c2_187  Protein name  Description  (EC 2.4.1)	NTID 758	<u>AAID</u> 2678	NT Length 397	pir:A53770  AA Score  I194 738  Locus Name  Sp:MURG_HAEIN	A53770  Probability  5.5e-73  Acc# P45065
growth factor-resp smooth muscle:SM-20 Description  ORF Name  16064061_c2_187  Protein name  Description  (EC 2.4.1)  ORF Name	NTID 758 NTID	AAID  AAID	NT Length 397 NT Length	pir:A53770  AA Score  Length 738  Locus Name  Sp:MURG_HAEIN  AA Score  Length Score	A53770  Probability  5.5e-73  Acc# P45065
growth factor-resp smooth muscle:SM-20 Description  ORF Name  16064061_c2_187  Protein name  Description  (EC 2.4.1)  ORF Name  16171905_c2_202	NTID 758 NTID	AAID  AAID	NT Length 397 NT Length	pir:A53770  AA Score Length 738  Locus Name Sp:MURG_HAEIN  AA Score Length Score 225	Probability  5.5e-73  Acc# P45065  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
16585933_t1_11	760	2680	765	2298	2390	4.8e-248
Protein name					s Name H_AZOVI	Acc# P16100
Description				<del></del>		<del>-</del>
DECARBOXYLASE) (I	DH)					
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
16678186_c3_251	761	2681	488	1467	1057	8.6e-107
Protein name				_	s Name	Acc#
hypothetical prot	ein F32D8	3.4		pir:T	21659	T21659
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20080051_c1_171	762	2682	299	900	330	9.4e-30
Protein name				Locu	s Name	Acc#
				sp:YJ	JA_ECOLI	P39408:P78
Description						143
HYPOTHETICAL 28.9	KD PROT	ETN TN OS	MY-DEOC IN	TERGENIC	REGION	
			NT	AA		
ORF Name	NTID	AAID	<u>Length</u>	Length	Score	Probability
20509628_c2_204	763	2683	378	1137	891	3.4e-89
Protein name					s Name	Acc#
				sp:FT	SZ_ECOLI	P06138:P78
Description						047:P77857
CELL DIVISION PRO	TEIN FTS	Z		-		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2132006_c2_218	764	2684	132	399	153	2.4e-10
Protein name				Locu	s Name	Acc#
hypothetical prot	ein sIII	330		pir:S	75232	S75232
Description			•	<del></del>		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22048442_f2_95	765	2685	402	1209	1010	8.2e-102
Protein name					s Name PE_HAEIN	Acc# P44514
Description					_	
SUCCINYL-DIAMINO	PIMELATE	DESUCCINY	LASE, (SDA	(P)	.,	-
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22323956_f3_130	766	2686	61	186	109	6.5e-06
Protein name				Locu	s Name	Acc#
hypothetical prot	ein PH022	21		pir:D	71245	D71245
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22463311_f3_128	767	2687	103	312		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22734807_f1_40	768	2688	97	294	,	
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22930306_c1_170	769	2689	354	1065	427	5.0e-40
Protein name				Locu	s Name	Acc#
5'-nucleotidase				gp:CL	1131243	AJ131243
Description						
Columba livia mRN	NA for 5'	-nucleoti	dase.		_	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23445136_£2_60	770	2690	583	1752	752	4.3e-120
Protein name NH(3)-dependent NAD	(+) syn	thetase	-11	<u>Locus</u> pir:G	S Name 72277	<u>Acc#</u> G72277
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
23532300_c1_165	771	2691	78	237		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT				***************************************		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23556625_c3_245	772	2692	241	726	185	2.2e-14
Protein name					S Name SQ_ECOLI	Acc# P06136
Description						
CELL DIVISION PROTE	IN FTSQ					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23615832_f1_16	773	2693	334	1005	125	9.1e-09
Protein name				Locus	s Name	Acc#
lysophospholipase he	omolog			pir:T	02661	T02661
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24111015_c3_269	774	2694	154	465	274	8.1e-24
Protein name					S Name BF_ECOLI	Acc# P08391:P75
Description						621

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
24219056_c1_184	775	2695	246	741	512	4.9e-49
Protein name				Locus sp:YPT5		Acc# P24562
Description				<u> </u>		
HYPOTHETICAL 24.5 K	D PROTE	IN IN PIL	r 5'REGIC	N (ORF5)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
24220786_f1_4	776	2696	370	1113	835	2.9e-83
Protein name				Locus sp:PILT		Acc# P24559
Description				<u>.                                    </u>		
TWITCHING MOBILITY	PROTEIN	Г				
ORF Name	NTID	AAID	NT Length	AA Length	core	Probability
24250928_c1_153	777	2697	78	237	207	1.0e-16
Protein name				Locus sp:YFHJ		Acc# P37096
Description				Sp. IIIIo		137030
HYPOTHETICAL 7.7 KL	PROTEI	N IN PPEB	-FDX INTE	RGENIC REG	ION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
24255260_c2_229	778	2698	115	348		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
24412562_c2_189	779	2699	314	945	740	3.4e-73
Protein name				Locus sp:DDL		Acc# P44405
Description					•	
D-ALANINED-ALANIN	E LIGAS	E, (D-ALA	NYLALANIN	E SYNTHETA	SE)	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
24415875_c1_174	780	2700	322	969	832	6.0e-	83
Protein name					s Name MZ_ECOLI		Acc# P23871:P78
Description							232
SYNTHETASE)			_				,
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score		bility
24644577_£2_96	781	2701	535	1608	923	1.4e-	92
Protein name					s Name		Acc#
hypothetical prote	1n			pir:S	76051		S76051
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
24813161_f3_143	732	2702	344	1035	650	1.2e-	63
Protein name				Locu	s Name		Acc#
MsmX				gp:AB	013374		AB013374
Description				-			
Bacillus haloduran complete cds.	s C-125	mamX, ујс	IA, ykoK a	nd yvfK	genes, pa	irtiala	nd
ORF Name	NTID	<u> </u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	bility
25579510_f3_109	783	2703	156	471	99	0.001	8
Protein name			•		s Name		Acc#
myosin alpha heavy	chain,	masticato	ry muscle	pir:S	33732		S33732
<u>Description</u>							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
26212750_c2_205	784	2704	329	990	285	7.8e-2	25
Protein name					s Name AC006436		<u>Acc#</u> AC006436
<u>Description</u>				<u> </u>			
Arabidopsis thalia sequence.	na chron	mosome II	BAC F13J1	1 genomic	c sequenc	e,comp	lete

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
26678567_c1_164	785	2705	63	192	88	0.00042
Protein name hypothetical protei	n 29.1			Locus pir:S	s <u>Name</u> 59084	Acc# S59084
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
26813135_c2_192	786	2706	524	1575	1287	3.7e-131
Protein name alkyl hydroperoxide	roduat	-200 F52A	profein		s Name	Acc#
	Teduce	.ase, roza	procein	pir:D	64 / 94	D64794
Description						
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
273427_c3_263	787	2707	224	675	583	1.5e-56
Protein name					s <u>Name</u> DA_ECOLI	Acc# P09548
Description				<u> </u>		
DEDA PROTEIN (DSG-1	PROTE	IN)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29337825_f2_62	788	2708	67	204	131	1.2e-08
Protein name					s Name T1_PSEAE	Acc# P24560
Description						
HYPOTHETICAL 17.0	D PROTE	EIN IN PIL	T 5'REGIO	N (ORF1)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31252614_c3_231	789	2709	68	207	85	0.0054
Protein name				Locus	s Name	Acc#
glutathione synthet	ase			gp:D8	8540	D88540
Description				<del></del>		
Synechococcus sp. I	ONA for	glutathio	ne synthe	tase, co	mplete co	is.

ORF Name	NTID A	AID	<u>NT</u> Length	AA Length	Score	Probability
3236505_c2_217	790	2710	119	360	127	1.7e-07
Protein name hypothetical prote	in sll1830			Locus pir:S	Name 75232	Acc# S75232
Description						
ORF Name		AID	NT Length	AA Length	Score	Probability
32593750_£1_27	791	2711	366	1101	1382	3.1e-141
Protein name					Name CA_ACICA	<u>Acc#</u> P42438
Description						
RECA PROTEIN						
ORF Name	NTID A	AID	NT Length	AA Length	Score	Probability
33788286_c2_188	792	2712	493	1482	1448	3.2e-148
Protein name UDP-N-acetylmuramat	e:L-alanii	ne ligas	e MurC	Locus	Name 110740	Acc# AF110740
Description		<u>.</u>	-	J [		
Pseudomonas aerugingene, complete cds.	nosa UDP-N	-acetyIm	uramate:	L-alanine	ligase	MurC(murC)
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34040777_c1_169	793	2713	196	591	93	0.00011
Protein name					Name DD_ECOLI	Acc# P36647
Description						
PREPILIN PEPTIDASE	DEPENDENT	PROTEIN	D PRECUI	RSOR		
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34159412_f1_53	794	2714	330	993	497	1.9e-47
Protein name oxidative stress to	ranscriptio	onal reg	ulator	Locus gp:XCU	Name J94336	<u>Acc#</u> U94336
Description						
Xanthomonas campes (ahpF) and oxidative cds.						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35276891_±3_132	795	2715	80	243	88	0.00042
Protein name hypothetical protei Description	n			Locu pir:D	s <u>Name</u> 75542	Acc# D75542
ORF Name 3907311_f3_110 Protein name	<u>NTID</u>   796	AAID 2716	NT Length 74	AA Length 225 Locu	Score	Probability  Acc#
Description						
NO-HIT						
ORF Name  3928750_c3_242  Protein name	<u>NTID</u> 797	AAID 2717	<u>NT</u> <u>Length</u> 298		Score 412 s Name	Probability  1.9e-38  Acc#
Description				Sp:In	IR_HAEIN	P31777
HYPOTHETICAL PROTEI	N HIO44	(ORFJ)				
ORF Name 3940943_t2_59	<u>NTID</u> 798	<u>AAID</u>	NT Length	AA Length 357	Score 93	Probability 0.00012
Protein name					S Name FE_ECOLI	Acc# P45580
Description						
HYPOTHETICAL 12.6 F	D PROTI	EIN IN PEP	P-SSR INT	ERGENIC	REGION (C	109)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3947318_f3_116	799	2719	276	831	1038	8.9e-105
Protein name					Name PD_MYCBO	Acc# P56220
Description						
(THP SUCCINYLTRANSE	ERASE)	(TETRAHYD)	ROPICOLIN	ATE SUCC	INYLASE)	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4017832_£2_85	800	2720	211	636	525	2.0e-50
Protein name  DedA family protein				Locus pir:B	8 Name 75253	Acc# B75253
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
4023342_c1_186	801	2721	215	648	145	3.8e-10
Protein name					Name FB_ECOLI	<u>Acc#</u> P25533
Description						
(F194)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
402336_f2_89	802	2722	83	252		
<u>Protein name</u>				Locus	Name	Acc#
Description						
NO-HIT					,	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4140943_t2_73	803	2723	157	474		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4331430_f1_28	804	2724	313	942	124	2.4e-16
Protein name					Name	Acc#
				sp:REC	CX_VIBCH	Q56647
Description						
REGULATORY PROTEIN	RECX					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4332837_c3_256	805	2725	105	318	149	6.7e-10
Protein name hypothetical protei	n sll18:	30		Locus	s Name 75232	Acc# S75232
<u>Description</u>		*.	<del>_</del>	ــــــــــــــــــــــــــــــــــــــ		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4348813_c3_259	806	2726	350	1053	1035	1.8e-104
<u>Protein name</u>					s <u>Name</u> P_HAEIN	Acc# P43764
Description						
(GLYCOPROTEASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4694427_f3_114	807	2727	362	1089	884	1.9e-88
Protein name					Name	Acc#
Description				sp:111	PA_HAEIN	P44463
LIPOIC ACID SYNTHET	ASE (LI	P-SYN) (L	IPOATE SY	NTHASE)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4773260_f2_79	808	2728	79	240		
Protein name				Locus	s Name	Acc#
Description						
мо-ніт						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4798536_c2_191	809	2729	212	639	685	2.3e-67
Protein name alkyl hydroperoxide	reducta	ase subuni	lt C		Name 129406	Acc# AF129406
Description				- <u> </u>		
Bacteroides fragili hydroperoxide reduct	_					(ahpC) and alkyl

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	ability
4824062_c2_200	810	2730	273	822	353	3.4e-	-32
Protein name					S Name		Acc#
Description							Q48269:007 681
(PHOSPHATIDYLSERIN	SYNTHA	SE)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	ability
6110943_c2_203	811	2731	435	1308	303	3.5e-	-26
Protein name	- `				SA_BUCAP	1	<u>Acc#</u> 051928
Description				<u> </u>			
CELL DIVISION PROTI	EIN FTSA						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	ability
659686_c2_210	812	2732	278	837	485	3.5e-	·46
Protein name					s Name		Acc#
Description				<b>5P</b> .10.			Q57097:005 009
HYPOTHETICAL PROTE	IN HIO11	8					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	ability
6754081_c3_235	813	2733	254	765	169	1.1e-	12
Protein name				Locus	s Name		Acc#
hypothetical prote	n MTH93	9		pir:G	69225		G69225
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	ability
681453_f3_127	814	2734	84	255	76	0.038	3
Protein name				-	s Name EH BACSU	1	<u>Acc#</u> P54947
Description				<u> </u>			~~~~
HYPOTHETICAL 30.2	KD PROTE	IN IN IDH	-DEOR INT	ERGENIC	REGION		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
682641_f1_42	815	2735	86	261	100	2.2e-05
Protein name hypothetical protei	n PH021	7		Locus	s <u>Name</u> 71244	Acc# G71244
Description			<u> </u>			
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
7222187_f1_35	816	2736	245	738	287	3.4e-25
Protein name conserved hypotheti	cal pro	tein ykrA		Locus pir:C	8 Name 69862	Acc# C69862
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
781563_c2_227	817	2737	113	342	265	7.3e-23
Protein name					s Name T6_PSEAE	Acc# P24564
Description						
HYPOTHETICAL 19.5 K	D PROTE	IN IN PIL	r region	(ORF6)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
812535_f1_43	818	2738	77	234		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10625252_f1_3	819	2739	581	1746	1790	1.8e-184
<u>Protein name</u>					S Name P_HAEIN	Acc# P43830
Description						
PROLYL-TRNA SYNTHET	ASE, (P	ROLINETI	RNA LIGAS	E) (PROR	5)	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
20496062_c2_30	820	2740	408	1227	1522	4.6e-1	.56
Protein name				******	Name PB_ACICA		<u>Acc#</u> P16706
Description							
TRYPTOPHAN SYNTHASE	BETA C	HAIN,			-		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
22847180_c1_17	821	2741	213	642	568	5.7e-5	55
Protein name					Name OG_ECOLI		<u>Acc#</u> P36879
Description							
HYPOTHETICAL ABC TR	ANSPORT	ER ATP-BI	NDING PRO	TEIN YADO	3		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
24642268_c3_37	822	2742	213	642	510	7.9e-4	9
Protein name					Name PF_ACICA		Acc# P16923
Description						_	
N-(5'-PHOSPHORIBOSY	L) ANTHR	ANILATE IS	SOMERASE,	(PRAI)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
24814125_c2_31	823	2743	285	858	541	4.1e-5	2
Protein name	a laba	aha: a			Name		Acc#
tryptophan synthase	alpna	Chain		gp:AF	L07094		AF107094
Description							
Rhodobacter sphaero cds; and tryptophan			-		-	_	ctial
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probab	ility
30727194_c2_24	824	2744	62	189			
<u>Protein name</u>				Locus	Name		Acc#
Description							
NO-HIT							

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
3941642_c2_25	825	2745	100	303 224	1.6e-18
Protein name		-		Locus Name	Acc# P36879
Description				•	
HYPOTHETICAL ABC	TRANSPOR	TER ATP-BI	NDING PRO	TEIN YADG	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4181557_c1_19	826	2746	281	846 553	2.2e-53
Protein name				Locus Name sp:YQCD_ECOLI	Acc# Q46920
Description				<u> </u>	
HYPOTHETICAL 32.6	KD PROT	EIN IN SYD	-SDAC INT	ERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4426338_c2_26	827	2747	260	783 744	1.3e-73
Protein name				Locus Name sp:YADH ECOLI	Acc#
Description				<u> </u>	P36880:P75 657
HYPOTHETICAL 28.5	KD PROT	EIN IN HPT	-PAND INT	ERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5120412_c3_32	828	2748	186	561 196	1.5e-15
Protein name				Locus Name	Acc#
cytochrome c5				gp:AVU94420	U94420
Description		•			
Azotobacter vinel cytochrome c5 (cyc xanthinephosphorik	B) gene,	complete	cds, and		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
7031312_c3_33	829	2749	66	201 129	7.5e-08
Protein name				Locus Name sp:YADG_ECOLI	<u>Acc#</u> P36879
Description				L	
HYPOTHETICAL ABC	TRANSPOR'	TER ATP-BI	NDING PRO	TEIN YADG	

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	pility
11198430_c3_60	830 2750	692	2079	3292	0.0	
Protein name  lactoferrin binding  Description	g protein B		Locus gp:AF(	Name 043131		Acc# AF043131
Moraxella catarrhal lactoferrin binding			_	_	_	
ORF Name 16128933 c2 53	NTID AAID	NT Length	AA Length [480	Score	Probab	oility
Protein name apolipoprotein N-ac	الــــا الــــا		Locus gp:AF0	Name 038595		Acc# AF038595
Description Pseudomonas aerugir cds.	nosa apolipoprote	-		se (cut	:)gene,	complete
ORF Name 19704378_c3_64	NTID AAID   [832   [2752 ]	NT Length	AA Length 1821	<u>Score</u> 1227	Probab 8.3e-1	
Protein name unknown			Locus	Name 043132		Acc# AF043132
Description			51 111			
Moraxella catarrhal lactoferrin binding					_	
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
24337826_c3_67	833 2753	718	2157	278	2.9e-2	21
Protein name hypothetical protei	n KOSH10 22			Name		Acc#
Description			pir:T2			T23512:T24

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	<u>Probability</u>
34164812_c2_54	834	2754	198	597	712	1.6e-80
Protein name  lactoferrin binding	protei	n B	, -		s Name 043133	Acc# AF043133
Description						
Moraxella catarrhal complete cds.	is stra	in VH19	lactoferri	n bindin	g protein	n B(lbpB) gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35837503_f2_26	835	2755	61	186		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35945257_c1_46	836	2756	67	204		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT			<del></del>			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3906263_c2_55	837	2757	1003	3012	5252	0.0
Protein name				Locus	s Name	Acc#
lactoferrin binding	protei	n A		gp:AF	043131	AF043131
Description						
Moraxella catarrhal lactoferrin binding						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3945388_£3_30	838	2758	413	1242	1219	5.9e-124
Protein name				Locus	s Name	Acc#
beta-ketoacyl-ACP s	ynthase	I		gp:PA	U70470	U70470
Description						
Pseudomonas aerugin gene, partial cds, b synthase I (fabB) ge	eta-hyd	roxy-ACP	dehydrase		_	-

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4005250_c3_68	839	2759	163	492	572	2.1e-55
ribosomal protein resistance protein	S12:stre	eptomycin		Locu pir:A	s Name 42939	Acc# B42939:A42 939:H64078
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4093767_c3_63	840	2760	544	1635	2854	3.2e-297
Protein name				Locus	s Name	Acc#
unknown				gp:AF	043131	AF043131
Description						
Moraxella catarrha lactoferrin binding						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4804632_f2_17	841	2761	485	1458	572	2.1e-55
		_				
Protein name	<b>_</b>	J [			S Name BB_SALTY	Acc# P12680
Protein name  Description		J I,				<del></del>
	E SYNTHAS	SE COMPONE	NT I, (AD	sp:PA	BB_SALTY	<del></del>
Description PARA-AMINOBENZOATE ORF Name	E SYNTHAS	SE COMPONE	NT I, (AL NT Length	sp:PA	BB_SALTY	<del></del>
Description PARA-AMINOBENZOATE	<u> </u>		NT	Sp:PA	BB_SALTY	P12680
Description PARA-AMINOBENZOATE ORF Name	NTID	AAID	<u>NT</u> Length	SP:PA	SE)	P12680  Probability
Description  PARA-AMINOBENZOATE  ORF Name  1018_c1_12	NTID	AAID	<u>NT</u> Length	SP:PA	SE) Score 163 Name	P12680  Probability  4.7e-12  Acc#
Description  PARA-AMINOBENZOATE  ORF Name  1018_c1_12  Protein name	NTID 842	AAID 2762	NT Length 229	Sp:PA	SE) Score 163 Name	P12680  Probability  4.7e-12  Acc#
Description  PARA-AMINOBENZOATE  ORF Name  1018_c1_12  Protein name  Description  Bacteriophage MB78  ORF Name	NTID 842	AAID 2762	NT Length 229 p26 & p2 NT Length	Sp:PA	SE) Score 163 Name	P12680  Probability  4.7e-12  Acc#
Description  PARA-AMINOBENZOATE  ORF Name  1018_c1_12  Protein name  Description  Bacteriophage MB78	NTID 842	AAID 2762 21, p11.5,	NT Length 229 p26 & p2	Sp:PA	SE) Score 163 S Name MB78P21	P12680  Probability  4.7e-12  Acc# X87092
Description  PARA-AMINOBENZOATE  ORF Name  1018_c1_12  Protein name  Description  Bacteriophage MB78  ORF Name	NTID  842  ORFs p2	AAID 2762 21, p11.5, AAID	NT Length 229 p26 & p2 NT Length	Sp:PA	SE) Score 163 S Name MB78P21	P12680  Probability  4.7e-12  Acc# X87092
Description  PARA-AMINOBENZOATE  ORF Name  1018_c1_12  Protein name  Description  Bacteriophage MB78  ORF Name  12303577_c1_11	NTID  842  ORFs p2	AAID 2762 21, p11.5, AAID	NT Length 229 p26 & p2 NT Length	Sp:PA	SE) Score 163 S Name MB78P21 Score	Probability  4.7e-12  Acc# X87092  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19557893_c2_17	844	2764	126	381		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT					<del></del>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20038305_c2_15	845	2765	75	228		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2134555_c1_13	846	2766	169	510		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23470625_c3_19	847	2767	189	570	287	5.9e-25
Protein name				· ·	s Name	Acc#
Description				gp:RP	4TRANOKF	L10330
Plasmid RP4 traN go complete cds.	ene, com	mplete cds	; traO ge	ne, comp	lete cds;	kirAgene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23632818_c3_25	848	2768	97	291		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
31555_c3_20	849	2769	333	1002 225	2.0e-17
Protein name				Locus Name	Acc#
Description	-				S58142:T42 283
ORF Name 34381286_c3_21	NTID 850	<u>AAID</u>	NT Length	AA Length 423	Probability
Protein name Description				Locus Name	Acc#
NO-HIT					•
ORF Name 34485637_c3_22	NTID 851	AAID 2771	NT Length	AA Length Score	Probability
Protein name Description				Locus Name	Acc#
NO-HIT					
ORF Name	NTID	AAID	NT Length	Length Score	Probability
Protein name Description	852	2772	81	Locus Name	Acc#
NO-HIT					
ORF Name [4804763_c2_16	NTID 853	<u>AAID</u>	<u>NT</u> Length	AA Length Score	Probability
Protein name Description NO-HIT			] [***]	Locus Name	<u>Acc#</u>
F					

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
6928168_c2_18	854	2774	308	927	
Protein name				Locus Name	Acc#
Description					
NO-HIT	· A.				
ORF Name	NTID	AAID	NT Length	AA Score	Probability
12625177_±1_2	855	2775	148	447 435	7.0e-41
Protein name				Locus Name sp:DKSA_ECOI	Acc# P18274
Description					
DNAK SUPPRESSOR PRO	TEIN		<del></del>		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14241635_c2_80	856	2776	130	393	
Protein name				Locus Name	Acc#
Description		•			
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
14876881_c2_82	857	2777	256	771 350	7.2e-32
Protein name				Locus Name	Acc# NO P17419
Description					
POSSIBLE FIMBRIAL A	SSEMBL	Y PROTEIN	FIMC (SER	OGROUP H1)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16181301_±3_63	858	2778	203	612 126	2.6e-06
Protein name				Locus Name sp:YGGH_ECOI	Acc# P32049
Description				L	
HYPOTHETICAL 27.3 I	CD PROT	EIN IN ANS	B-MUTY IN	TERGENIC REGION	(F239)

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
16898413_c1_77	859	2779	100	303	286	9.1e-24
Protein name					s Name 12_HAEIN	Acc# P44836
Description						
PROBABLE TONB-DEPEN	NDENT RI	ECEPTOR HI	0712 PREC	URSOR		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21517252_f1_3	860	2780	621	1866	1417	6.1e-145
Protein name					s Name RC_PSEFL	Acc# P32966
Description						· ·
EXCINUCLEASE ABC SU	BUNIT		-			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
22004587_f2_24	861	2781	333	1002	436	5.5e-41
Protein name				Locus	s Name	Acc#
Description				sp:YA	DB_ECOLI	P27305:P75 662
HYPOTHETICAL 34.9 K	D PROTE	EIN IN PCN	B-DKSA IN	TERGENIC	REGION	
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
23597187_±3_61	862	2782	769	2310	539	3.0e-100
<u>Protein name</u>					Name	Acc#
				sp:PR	IA_RHORU	P05445
Description						
PRIMOSOMAL PROTEIN	N' (RE	PLICATION	FACTOR Y)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23865681_f3_58	863	2783	739	2220	789	2.2e-78
<u>Protein name</u>					Name OT_HAEIN	Acc# P43811
Description						
((PPGPP)ASE) (PENTA	-PHOSPE	ATE GUANO	SINE-3'-P	YROPHOSPH	OHYDROLA	SE)

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24412817_c2_92	864	2784	84	255		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2510887_c1_76	865	2785	60	183		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT			<u>,</u> ,			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26594686_f1_14	866	2786	215	648	355	2.1e-32
Protein name					s Name CG_ECOLI	Acc#
Description						P31432:P76 720
HYPOTHETICAL 22.0	KD PROTE	IN IN RPH	-GMK INTE	RGENIC R	EGION PRI	ECURSOR
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29304827_c2_94	867	2787	63	192	286	9.1e-24
Protein name					s Name 12_HAEIN	<u>Acc#</u> P44836
Description				<u> </u>		<u></u>
PROBABLE TONB-DEPE	NDENT RE	CEPTOR HI	0712 PREC	URSOR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
29484418_c2_95	868	2788	566	1701	710	1.6e-118
Protein name					s Name	Acc#
methyltransferase				gp:AF	060119	AF060119
Description						
Pasteurella haemol (res) genes, comple		ethyltrans	ferase (π	nod) and	restrict:	ionendonuclease

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
30281911_c3_114	869 2789	79	240 286	9.1e-24
Protein name			Locus Name sp:Y712_HAEIN	Acc# P44836
Description				
PROBABLE TONB-DEPEN	NDENT RECEPTOR HIC	712 PREC	URSOR	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
34017010_£2_35	870 2790	307	924 708	8.3e-70
Protein name			Locus Name	Acc#
hypothetical protei	n b2431		pir:F65017	F65017
Description		<del>-</del>	<u> </u>	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
34407905_c2_93	871 2791	460	1383 1801	1.2e-185
Protein name			Locus Name	Acc#
L-2,4-diaminobutyra	te:2-ketoglutarat	.e	gp:AB001599	AB001599
Description				<del></del>
Acinetobacter bauma 4-aminotransferase,		-diamino	butyrate:2-ketog	Lutarate
ORF Name	NTID AAID	<u>NT</u> Length	AA Score	Probability
34641550_c3_119	872 2792	186	561 126	1.2e-07
Protein name			Locus Name sp:AIL_YEREN	Acc# P16454
Description			1	
ATTACHMENT INVASION	LOCUS PROTEIN PR	ECURSOR	····	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
3923818_f2_34	873 2793	62	189	
Protein name			Locus Name	Acc#
Description				
NO-HIT				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4306455_c1_65	874	2794	323	972	122	4.9e-07
Protein name					s Name P1_PSEAE	Acc# P17838
Description						
FIMBRIAL PROTEIN P	RECURSOR	(PILIN)	(STRAIN P	(1)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4329518_±3_56	875	2795	212	639	541	4.1e-52
Protein name					s <u>Name</u> UA_ECOLI	Acc# P24234
Description						
GUANYLATE KINASE,	(GMP KIN	ASE)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4428413_c1_78	876	2796	517	1554	2128	2.8e-220
Protein name					s Name	Acc#
L-2,4-diaminobutyra	ite deca:	rboxylase		gp:AC	CL24DD	D55724
Description						
Acinetobacter bauma complete cds.	annii ge	ne for L-	2,4-diami	nobutyra	tedecarbo	oxylase,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4538558_f3_51	877	2797	270	813	611	1.6e-59
Protein name					s Name	Acc#
hypothetical prote	<u> </u>			gp:PP	PAL1	X74218
Description						
Pseudomonas putida	ruvB, t	olQ, tolR	, tolA, t	olB and	oprL gene	es.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4570318_£3_44	878	2798	228	687	448	3.0e-42
Protein name					s <u>Name</u> H_HAEIN	Acc# P44755
Description						
PHOSPHOGLYCOLATE PI	ЮЅРНАТА	SE, (PGP)				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4902305_c2_96	879	2799	869	2610	2182	5.3e-226
Protein name restriction endonuc	lease				S Name 060119	Acc# AF060119
Description					·	
Pasteurella haemoly (res) genes, complet		thyltrans	terase (m	od) and i	restricti	onendonuclease
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4964000_f3_57	880	2800	95	288	187	1.3e-14
Protein name					S Name DZ_HAEIN	Acc# P43740
Description						
OMEGA CHAIN) (RNA P	OLYMERA	SE OMEGA	SUBUNIT)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
5079408_f3_60	881	2801	151	456	367	1.1e-33
Protein name hypothetical protei	n 1 (vn	fA 5' regi	ion)	Locus pir:B	Name 14514	Acc# B44514
Description				<u> </u>		
ORF Name 5080293_c2_89	NTID 882	AAID   2802	NT Length 592	AA Length	Score	Probability  8.5e-116
Protein name					Name	Acc# P45112
Description						
SINGLE-STRANDED-DNA	-SPECIF	'IC EXONUC	LEASE REC	J,		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
5339762_c3_109	883	2803	325	978	1184	3.0e-120
Protein name				_	Name 027189	Acc# AF027189
Description						<del></del>
Acinetobacter sp. B and unknown genes.	D413 ly	tB, comB,	comC, co	mE, and c	comF gene	s, complete cds;

ORF Name	NTID AAID	NT Length	AA Length	Probability
97582_f3_42	884 2804	289	870 291	1.3e-25
Protein name			Locus Name sp:ICC_ECOLI	Acc# P36650
Description				<del></del>
ICC PROTEIN				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
13673437_c2_39	885 2805	597	1791 150	3.5e-13
Protein name			Locus Name	Acc#
putative terminase			gp:AF147978	AF147978
Description	•			
Bacteriophage D3 pu protease, and major				
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
14181292_c3_43	2806	101	306	
Protein name			Locus Name	Acc#
Description				
по-ніт				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
19953125_c2_36	887 2807	215	648	
Protein name			Locus Name	Acc#
Description				
NO-HIT				
ORF Name	NTID AAID	<u>NT</u> Length	AA Score	Probability
25665952_c1_33	888 2808	128	387 88	0.0036
Protein name			Locus Name  gp:BPH251805	Acc#
Description			] Ab.peu521002	AJ251805
Bacteriophage phi-N	ZeO3-12 complet	e genome.		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
26675311_c2_37	889	2809	107	324	145	3.8e-1	0
Protein name hypothetical protei	n				S Name E133022		<u>Acc#</u> AJ133022
Description							
Xenorhabdus nemator	hilus p	roviral O	RF1 to OR	F8.			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	
32657262_c2_35	890	2810	150	453	209	6.7e-1	8
Protein name  DNA primase				Locus pir:C	<u>Name</u>		<u>Acc#</u> C41830
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
3941887_c1_32	891	2811	185	558	88	0.019	
Protein name				Locus gp:PF	Name		<u>Acc#</u> X17490
Description							
Plasmodium falcipar	cum mRNA	for aspa	ragine-ri	ch antig	en (clone	e53C6).	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	
5916253_c1_28	892	2812	257	774	259	3.1e-2	2
Protein name					s Name 22_HAEIN		Acc# P44193
Description							
HYPOTHETICAL PROTE	IN HI142	2					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	ility
900215_c3_41	893	2813	797	2394	330	4.7e-2	7
Protein name putative DNA primas	se				s Name 139719		<u>Acc#</u> AF139719
Description		· · ·		<u> </u>			
Klebsiella oxytoca cds; and unknown ger	_	pACM1 pu	tative DN	A primas	e (pri) g	gene, com	plete

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probab	oility
10959627_f2_14	894	2814	294	885	513	3.8e-4	19
Protein name				-	s Name EX_ECOLI		<u>Acc#</u> P77392
Description							
HYPOTHETICAL 33.3 K	D PROTE	IN IN CUT	E-ASNB IN	TERGENIC	REGION		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
1178127_f3_20	895	2815	963	2892	2605	1.9e-2	281
Protein name				Locus	s Name		Acc#
SecA				gp:AB	012226		AB012226
Description						•	
Vibrio alginolyticu	s gene	for SecA,	complete	cds.			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probak	oility
12298468_f1_13	896	2816	102	309	82	0.017	
Protein name				Locus	s Name		Acc#
probable membrane p	rotein	L549.12		pir:T	02800		T02800
Description				`			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
12985393_f3_16	897	2817	270	813	537	1.le-5	51
Protein name					s Name PD HAEIN		<u>Acc#</u> P44817
Description					_		
(PEPTIDASE D)			-				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
14538262_c2_40	898	2818	69	210	109	2.5e-0	06
Protein name				Locus	s Name		Acc#
hypothetical protei	n APE04	58		pir:A	72741		A72741
Description			<u> </u>				

ORF Name	NTID	AAID	<u>NT</u> Length	Length Sco	ore	Probability
14548260_c1_34	899	2819	244	735	65	7.3e-23
Protein name hypothetical pro	tein D1022	2.4		Locus Na		Acc# T34190
Description						
ORF Name	NTID	AAID	NT Length	Length	ore	Probability
21640900_f1_8	900	2820	442	1329	361	5.3e-139
Protein name				Locus Na		Acc# P48247
Description						
(GLUTAMATE-1-SEM	TALDEHYDE	AMINOTRA	NSFERASE)	(GSA-AT)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
24609561_c3_56	901	2821	241	726 1	47	2.4e-19
Protein name				Locus Na		Acc#
Description				sp:UP14_	ECOLI	P39179:Q46 826
UNKNOWN PROTEIN	FROM 2D-PA	AGE (SPOT	PR51)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
36329806_f1_2	902	2822	80	243 6	0	0.019
Protein name	- Fabrar	-Founda	22	Locus Na		Acc#
thyroid hormone Description	<u>surrotrans</u>	ierase, i		pir:JC58	35	JC5885
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
4188811_±1_6	903	2823	159		10	1.2e-27
4188811_f1_6  Protein name  conserved hypoth	903				<u>ıme</u>	1.2e-27 Acc# T03501

ORF Name	NTID 2	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
4336018_f1_7	904	2824	115	348	349	9.1e-3	32
Protein name					S Name NA_ECOLI		Acc# P16680
Description							
PHNA PROTEIN							
ORF Name	NTID A	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
4689143_f2_15	905	2825	175	528	132	9.0e-0	8
Protein name apolipoprotein N-ac	cyltransfe	erase		_	Name 038595		Acc# AF038595
Description				J [21			
Pseudomonas aerugi cds.	nosa apoli	poprotei	n N-acyl	transfera	ase (cut	) gene,	complete
ORF Name	NTID A	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
5205312_f3_21	906	2826	69	210			
Protein name				Locus	Name		Acc#
Description							
NO-HIT							
ORF Name	NTID Z	AAID	NT Length	<u>AA</u> Length	Score	Probab	oility
22382752_c1_11	907	2827	117	354	100	2.2e-0	5
Protein name				Locus	Name		Acc#
hypothetical prote	in			pir:T	10511		T10511
Description		*					
ORF Name	NTID A	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
26251376_c1_8	908	2828	82	249			
Protein name				Locus	Name		Acc#
Description							
NO-HIT	-				<del></del>		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35188942_c3_14	909	2829	255	768	248	4.6e-21
Protein name hypothetical prote	ein slr1	971			s Name 75639	Acc# S75639
Description				<del></del>	. <del>-</del>	
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
35337805_c1_10	910	2830	185	558	152	5.9e-12
Protein name					s Name	Acc#
sulfate transporte	er ———			gp:D8	9631	D89631
Description						
Arabidopsis thali	ana mRNA	for sulf	ate transp	orter, c	omplete d	cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35801416_f1_1	911	2831	255	768	379	6.1e-35
Protein name					s Name	Acc#
				sp:RL	UA_ECOLI	P39219
Description				sp:RL	UA_ECOLI	P39219
Description (PSEUDOURIDYLATE	SYNTHASE	) (URACIL	HYDROLYAS	<u> </u>	UA_ECOLI	P39219
-	SYNTHASE NTID	) (URACIL	HYDROLYAS <u>NT</u> Length	<u> </u>	UA_ECOLI Score	P39219 Probability
(PSEUDOURIDYLATE	· · · · ·		NT	EE) <u>AA</u>		
(PSEUDOURIDYLATE	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
ORF Name 4328403_c2_13	NTID	AAID	<u>NT</u> Length	AA Length 330 Locu	Score	Probability 5.1e-15
ORF Name  4328403_c2_13  Protein name	NTID	AAID	<u>NT</u> Length	AA Length 330 Locu	Score 191 s Name	Probability  5.1e-15  Acc#
ORF Name  4328403_c2_13  Protein name  BolA protein	<u>NTID</u>	<u>AAID</u>	NT Length 109	AA Length 330 Locu gp:PF	Score 191 s Name L243174	Probability  5.1e-15  Acc# AJ243174
(PSEUDOURIDYLATE  ORF Name  4328403_c2_13  Protein name  BolA protein  Description	<u>NTID</u>	<u>AAID</u>	NT Length 109	AA Length 330 Locu gp:PF	Score 191 s Name L243174	Probability  5.1e-15  Acc# AJ243174
ORF Name  4328403_c2_13  Protein name  BolA protein  Description  Pseudomonas fluor	NTID 912 escens p	AAID  2832  artial Fu	NT Length 109 marase C g	AA Length 330 Locu gp:PF	Score  191  S Name  L243174  A gene ar	Probability  5.1e-15  Acc# AJ243174  ndORF1.
ORF Name  4328403_c2_13  Protein name  BolA protein  Description  Pseudomonas fluor  ORF Name	NTID  912  escens p	AAID  AAID  AAID	Length  109  marase C g  NT Length	AA Length  AA Length  AA Length  AA Length	Score  191  S Name  L243174  A gene ar  Score	Probability  5.1e-15  Acc# AJ243174  ndORF1.  Probability
ORF Name  4328403_c2_13  Protein name  BolA protein  Description  Pseudomonas fluor  ORF Name  4572206_c1_9	NTID  912  escens p  NTID  913	AAID  AAID  AAID	Length  109  marase C g  NT Length	AA Length  330  Locu  gp:PF  gene, bol.  AA Length  1143  Locu	Score  191 s Name L243174 A gene ar Score 391	Probability  5.1e-15  Acc# AJ243174  ndORF1.  Probability  1.5e-35
ORF Name  4328403_c2_13  Protein name  BolA protein  Description  Pseudomonas fluor  ORF Name  4572206_c1_9  Protein name	NTID  912  escens p  NTID  913	AAID  AAID  AAID	Length  109  marase C g  NT Length	AA Length  330  Locu  gp:PF  gene, bol.  AA Length  1143  Locu	Score  191 s Name L243174 A gene ar Score 391 s Name	Probability  5.1e-15  Acc# AJ243174  adoRF1.  Probability  1.5e-35  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16990667_±1_7	914	2834	61	186	109	8.8e-06
Protein name hemV protein			**	Locus pir:S	Name 54440	Acc# S54440
Description						
ORF Name	NTID   [915	<u>AAID</u>	<u>NT</u> <u>Length</u>	AA Length 213	Score	Probability
Protein name	] [913		] [/"		s Name	Acc#
Description					***************************************	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19573425_c1_31	916	2836	150	453		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20408375_c2_34	917	2837	60	183		
Protein name				Locus	Name	Acc#
Description						
NO-HIT		· · · · · · · · · · · · · · · · · · ·				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20930_c3_42	918	2838	144	435	87	0.0070
Protein name					Name PZ PSEPU	Acc# P25760
Description						
ATP SYNTHASE PROTE	IN I					

ORF Name	NTID AAID NT Length	AA Length <u>Score</u>	Probability
20991307_c2_35	919 2839 302	909 128	8.0e-14
Protein name periplasmic zinc tr Description	ansporter ZnuA	Locus Name	Acc# AF141971
and periplasmic zinc	HI0318 homolog gene, pa transporter ZnuA (znuA) somerase A homologgene,	genes, complete co	
ORF Name	NTID AAID NT Length	AA Score Length 1866	Probability
Protein name	synthase, beta chain	Locus Name pir:D64071	Acc# D64071
Description	NT	7.7	
ORF Name [24507777 c3 43	NTID AAID Length  [921 [2841 [520]]	AA Score Length 1563 1979	Probability
Protein name		Locus Name sp:ATPA_ECOLI	Acc# P00822
Description ATP SYNTHASE ALPHA	CHAIN,	10	
ORF Name 25680186_c1_28	NTID AAID NT Length  922 2842 160	<u>AA</u> <u>Length</u> Score  483  371	Probability 4.3e-34
<u>Protein name</u>		Locus Name sp:ATPF_VIBAL	Acc# P12989
Description  ATP SYNTHASE B CHAI	N		
AIP SININASE B CHAI	NT NT	7 A	
ORF Name 34023378 cl 26	NTID AAID Length    923   2843   296	<u>AA</u> <u>Length</u>   Score     745	Probability  9.9e-74
Protein name		Locus Name	Acc#
Description		sp:ATP6_ECOLI	P00855:Q47
ATP SYNTHASE A CHAI	N. (PROTEIN 6)		

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	<u>Probability</u>
35159406_c3_39	924	2844	65	198 65	0.0045
Protein name extensin homolog	F2401.18			Locus Name	Acc# T01456
Description					
ORF Name	NTID	AAID	NT Length	AA Score	Probability
3953587_f3_23	925	2845	174	525 236	8.6e-20
Protein name				Locus Name sp:ZUR_ECOLI	Acc# P32692:P76
Description					784
ZINC UPTAKE REGUL	ATTON DR	OTEIN (ZI	NC HPTAKE	PEGIII.ATOR)	
Zine or race		O1211 (21		THE PERSON NAMED IN	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4120425_c1_29	926	2846	202	609 268	3.5e-23
Protein name				Locus Name	Acc#
				sp:ATPD_VIBA	L P12987
Description					
ATP SYNTHASE DELT	A CHAIN,				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4332943_c1_27	927	2847	84	255 261	1.9e-22
Protein name				Locus Name sp:ATPL_HAEI	Acc# N P43721
Description					
(DICYCLOHEXYLCARE)	ODIIMIDE	-BINDING	PROTEIN)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
56333_c3_44	928	2848	309	930 894	1.6e-89
<u>Protein name</u>					
				Locus Name	Acc#
Description				Locus Name sp:ATPG_ECOL	

ORF Name	NTID AA	ID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
7056441_c2_38	929 2	849	139	417	300	1.4e-	26
Protein name					Name PE_HAEIN		Acc# P43718
Description							
ATP SYNTHASE EPSILO	ON CHAIN,						
ORF Name	NTID AA	ID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
9928433_t2_17	930 2	850	71	216			
Protein name				Locus	Name		Acc#
Description				•			
NO-HIT							
ORF Name	NTID AA	ID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
3235950_c2_28	931 2	851	561	1686	2096	6.8e-	217
Protein name				Locus	Name		Acc#
urocanase				gp:PSI	EHUTUU		M22002 M00
Description							M33923:M28 362
Pseudomonas putida	urocanase	(hutU)	gene, co	mplete co	is.		
ORF Name	NTID AA	ID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
3940938_c2_30	932 2	852	357	1074	366	1.4e-	33
Protein name					Name		<u>Acc#</u> P19452
Description				<u> </u>			1 101
(HISTIDINE UTILIZAT	'ION PROTEI	N G) (F	RAGMENT)				
ORF Name	NTID AA	ID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
3953181_c3_34	933 2	853	434	1305	1007	1.7e-	101
Protein name					Name		Acc#
Description				gp:YP1			AL031866
Yersinia pestis 102	kbases un	stable	region:	from 1 to	119443.		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4822181_f3_14	934	2854	78	237		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT					<del></del>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
789037_c3_33	935	2855	525	1578	1486	3.0e-152
Protein name					s Name	Acc#
histidine ammonia-l	yase,:	nistidase		pir:A	35251	A35251:S39
Description						381
ORF Name 828942_t1_1	<u>NTID</u>	<u>AAID</u>	NT Length 296	AA Length 891	Score	Probability
Protein name	] [			Locus	S Name	Acc#
					AM_BACSU	P37511
Description						
HYPOTHETICAL 32.9 F	D PROT	EIN IN TET	B-EXOA IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10978400_c3_163	937	2857	253	762	186	1.7e-14
Protein name				-	Name M4_PSEAE	Acc# P48246
Description						
p	<del></del>					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1259626_c2_127	938	2858	614	1845	1756	7.3e-181
Protein name					Name	Acc#
Description				sp:YA	51_HAEIN	Q57180:005 043
HYPOTHETICAL ABC TR	ANSPOR'	TER ATP-BI	NDING PRO	TEIN HI10	051	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12672305_c2_140	939	2859	110	333	144	4.8e-10
<u>Protein name</u>					s Name GX_HAEIN	Acc# P44048
Description						
HYPOTHETICAL PROTE	IN HIO7	60				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13131890_£3_71	940	2860	169	510	147	2.7e-10
Protein name					s Name BC_ERWCH	Acc# P39691
Description						
THIOL:DISULFIDE IN	TERCHAN	GE PROTEIN	DSBC PRE	CURSOR		
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
14847290_£3_89	941	2861	61	186		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
15875832_f3_92	942	2862	75	228		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT					····	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16493891_c3_165	943	2863	118	357		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT					-	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16610052_c3_183	944	2864	93	282		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT				*****		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1957701_f1_30	945	2865	277	834	329	1.2e-29
Protein name					s Name PE_HAEIN	<u>Acc#</u> P43732
Description				<u>L</u>		
GRPE PROTEIN			<del>.</del>	<u> </u>	<del>.</del>	7
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19720930_f3_74	946	2866	276	831	597	4.8e-58
Protein name					s Name PB_ECOLI	Acc# P04036
Description						
DIHYDRODIPICOLINA	TE REDUC	rase,				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21489390_t2_59	947	2867	67	204		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT				<del>,</del>		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
21689063_±1_10	948	2868	414	1245	328	1.5e-29
Protein name				Locus	s Name	Acc#
SrpJ				gp:AF	176824	AF176824
Description						
Synechococcus PCC gamma-glutamyltram SrpF (srpF), SrpJ membrane lipoprote genes, complete co	nspeptidas (srpJ), A ein SrpL	se SrpE ( ATP-bindi	srpE), alp ng protein	ha-helica of ABC to	alcoiled- ransporte	coil protein er SrpK (srpK),

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22042177_f2_65	949	2869	493	1482	1203	2.9e-122
Protein name argininosuccinate I	yase arg	эн		Locus pir:C	s <u>Name</u> 69589	<u>Acc#</u> C69589
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22070191_f1_5	950	2870	409	1230	486	2.8e-46
Protein name				Locus	s Name	Acc#
cystathionine-gamma	-lyase			gp:AF	180145	AF180145
Description					***	
Zymomonas mobilis G protein yidC (yidC), gltB (gltB), glutama udk (udk), hypothetic protein; zml2orf5, hy A, beta-hydroxysteroi	hypoth te syntl al prote pothetic	etical pro hasesmall ein, NADH cal prote:	otein,glu subunit dehydrog in, aspar	tamine-py gltS (glt enase, hy tate amin	yruvate a tS), unde ypothetic notransfe	mminotransferase ecaprenol kinase cal
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
22922082_c2_138	951	2871	319	960		
Protein name				Locus	s_Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
234701_c3_187	952	2872	73	222	115	5.1e-07
Protein name				Locus	Name	Acc#
extensin				pir:S	22697	
Description						S22697:S21 006
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23572127_f1_31	953	2873	636	1911	2307	3.0e-239
Protein name					Name AK_FRATU	Acc# P48205
Description				<u> </u>		
DNAK PROTEIN (HEAT	SHOCK PI	ROTEIN 70)	(HSP70)	•		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23909688_±3_77	954	2874	466	1401	456	4.2e-43
Protein name rubredoxinNAD+ re protein hydA 3'-regi		,:hypothe	tical	Locus pir:C	Name 65051	Acc# C65051
Description				_		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24073762_c3_158	955	2875	220	663	681	6.0e-67
Protein name AvtA					Name 014804	Acc# AF014804
Description				J <u>2.                                    </u>		
Neisseria meningiti genes, complete cds.		B (pgIB),	PglC (pg	IC), Pgll	(pgID),	andAvtA (avtA)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
24100465_£2_46	956	2876	321	966	639	1.3e-64
Protein name intrinsic membrane	profein	<u>, , , , , , , , , , , , , , , , , , , </u>			Name	Acc# AB000100
Description	Process			J gp:Ab	300100	AB000100
Synechococcus sp. D cyanase, complete cd		intrinsic	membrane	protein	malK-li	keprotein,
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
24391877_c1_109	957	2877	329	990	927	5.1e-93
Protein name					Name	Acc#
Description				Sp. IIII	M3_ECOLI	P06983:P78 125
SYNTHASE) (HMBS) (P	RE-UROP	ORPHYRINO	GEN SYNTH	ASE)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24417807_f3_76	958	2878	185	558	263	1.2e-22
Protein name Mip				Locus	Name	Acc# S71704
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
24619003_£3_79	959	2879	287	864 625	5.2e-61
Protein name				Locus Name	Acc# P73450
Description					
NITRATE TRANSPORT	ATP-BIN	DING PROTE	IN NRTC		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
26053250_c1_112	960	2880	268	807	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
26053250_c2_134	961	2881	116	351	
Protein name				Locus Name	Acc#
Description					
NO-HIT				****	
110 1111					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
	NTID 962	AAID 2882		Score	Probability
ORF Name			Length	Ength 891 480  Locus Name	1.2e-45 <u>Acc#</u>
ORF Name 26362680_c2_129			Length	Length Score 480	1.2e-45 <u>Acc#</u>
ORF Name  26362680_c2_129  Protein name	962	2882	Length 296	Length 480  Locus Name  sp:YJFH_HAEI	1.2e-45 <u>Acc#</u>
ORF Name  26362680_c2_129  Protein name  Description	962	2882	Length 296	Length 480  Locus Name  sp:YJFH_HAEI	1.2e-45 <u>Acc#</u>
ORF Name  26362680_c2_129  Protein name  Description  HYPOTHETICAL TRNA	962 /RRNA ME	2882	Length 296 PERASE HIO	Length 891 480  Locus Name Sp:YJFH_HAE1  860,  AA Score	1.2e-45 Acc# N P44906
ORF Name  26362680_c2_129  Protein name  Description  HYPOTHETICAL TRNA,  ORF Name	962 /RRNA ME* NTID 963	2882 THYLTRANSF  AAID  2883	Length 296 ERASE HIO NT Length 323	Length Score    891   480     Locus Name     Sp:YJFH_HAEI     860,     AA     Length   Score	1.2e-45  Acc# P44906  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29339432_c3_166	964	2884	322	969	92	4.2e-06
Protein name hypothetical prote	in b275	5		Locu pir:G	s <u>Name</u> 65056	<u>Acc#</u> G65056
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
29962837_c3_181	965	2885	288	867	117	9.9e-13
Protein name				· · · · · · · · · · · · · · · · · · ·	s Name AJ_SYNP7	<u>Acc#</u> P50026
Description						
DNAJ PROTEIN				-		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
32629186_c2_139	966	2886	223	672		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3307_£2_47	967	2887	119	360	349	9.1e-32
Protein name					s Name	<u>Acc#</u>
Description				sp:YA	DR_HAEIN	P45344
HYPOTHETICAL PROTE	IN HI17	23				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34492161_c3_168	968	2888	353	1062		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3940925_c3_182	969	2889	341	1026	466	3.7e-44
Protein name					s Name ET_ECOLI	Acc# P45524
Description						
HYPOTHETICAL 38.5	KD PROT	EIN IN KI	FB-PRKB IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4095443_c3_170	970	2890	161	486	147	2.3e-10
Protein name hypothetical protein	ein Rv01	63		Locus pir:G	Name 70903	<u>Acc#</u> G70903
Description				J [		
ORF Name 4187538_f2_50	<u>NTID</u>	<u>AAID</u>	<u>NT</u> <u>Length</u>	AA Length 1422	<u>Score</u>	Probability 1.1e-115
Protein name				Locus	s Name	Acc#
Description		- Order		sp:MP	L_HAEIN	P43948
2101027			NT	7.7		
ORF Name	NTID	AAID	<u>Length</u>	AA Length	Score	Probability
4328135_f1_13	972	2892	474	1425	812	7.9e-81
Protein name				Locus	s Name	Acc#
periplasmic subst	rate bind	ding prote	ein	gp:AF	001333	AF001333
Description						
Synechococcus PCC membrane protein (						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4328965_f3_99	973	2893	132	399	282	1.2e-24
Protein name					Name	Acc# 030825
Description						
HYPOTHETICAL PROT	EIN HYPO	117				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4423318_c3_160	974	2894	626	1881	2307	3.0e-239
Protein name 93% identity over coli	631 amir	no acids w	ith E.	_	S Name YSTMF1	Acc# AF170176
Description						
Salmonella typhim	urium fr	agment STM	FI.			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4454512_±1_32	975	2895	111	336	113	9.3e-07
Protein name					Name 73_HAEIN	Acc# P43960
Description						
HYPOTHETICAL PROT	EIN HIOT	73				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score 、	Probability
4548188_f1_38	976	2896	421	1266	1342	5.4e-137
Protein name					Name DA_ECOLI	Acc# P29011
Description						
D-AMINO ACID DEHY	DROGENAS	E SMALL SU	BUNIT,			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
5119127_c2_153	977	2897	651	1956	1464	6.4e-150
Protein name					s Name	Acc#
				sp:YH	ES_ECOLI	P45535
Description						
HYPOTHETICAL ABC	TRANSPOR'	TER ATP-BI	NDING PRO	TEIN YHES	5	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6727086_£1_17	978	2898	563	1692	879	6.3e-88
Protein name putative gamma-gl	utamylcys	steine syn	thetase		Name P243941	Acc# AJ243941
Description				<u> </u>		
Pseudomonas sp. s	train HR	199 partia	l vanB, f	dh, gcs,	ehyA and	ehyBgenes.

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
673253_£3_72	979	2899	407	1224 1146	3.2e-116
Protein name				Locus Name sp:DNAJ_SALTY	Acc# Q60004
Description				<u> </u>	
DNAJ PROTEIN		···			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
6854677_c3_167	980	2900	462	1389	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
801452_f2_60	981	2901	76	231	
Protein name				Locus Name	Acc#
Protein name Description				Locus Name	Acc#
				Locus Name	Acc#
Description	NTID	AAID	<u>NT</u> Length	Locus Name  AA Length Score	Acc# Probability
Description NO-HIT	NTID	<u>AAID</u>	_	AA Score	
Description  NO-HIT  ORF Name  884712_c1_113  Protein name	982	2902	Length	AA Score Length 79 Locus Name	Probability 0.0031 Acc#
Description  NO-HIT  ORF Name  884712_c1_113  Protein name hypothetical protes	982	2902	Length	<u>AA</u> <u>Length</u> 3576  79	Probability 0.0031
Description  NO-HIT  ORF Name  884712_c1_113  Protein name	982	2902	Length 1191	AA Score Length 79 Locus Name pir:A71069	Probability 0.0031 Acc#
Description  NO-HIT  ORF Name  884712_c1_113  Protein name hypothetical protes	982	2902	Length	AA Score Length 79 Locus Name	Probability 0.0031 Acc#
Description  NO-HIT  ORF Name  884712_c1_113  Protein name hypothetical protes  Description	982	2902	Length 1191 NT	AA Score  Length  3576  Tocus Name  pir:A71069  AA Score	Probability  0.0031  Acc# A71069
Description  NO-HIT  ORF Name  884712_c1_113  Protein name hypothetical protes  Description  ORF Name	982 in PH124  NTID	2902 16 <u>AAID</u>	Length 1191 NT Length	AA Score Length 79 Locus Name pir:A71069  AA Score	Probability  0.0031  Acc# A71069
Description  NO-HIT  ORF Name  884712_c1_113  Protein name hypothetical prote: Description  ORF Name  915633_c1_115	982 in PH124  NTID	2902 16 <u>AAID</u>	Length 1191 NT Length	AA Score Length 79 Locus Name pir:A71069  AA Score Length Score 225	Probability  O.0031  Acc# A71069  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
9766888_±1_18	984	2904	182	549	229	4.8e-19
Protein name				1.1-4.7	s <u>Name</u> 21_PSEAE	Acc# P21482
Description				4_6		<del>.</del>
HYPOTHETICAL 17.8	KD PROTI	EIN IN ALG	R2 5'REGI	ON		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10329680_£1_3	985	2905	544	1635	587	5.5e-57
Protein name				Locus gp:PS	S Name EOPRC	Acc# D28119
Description						
Pseudomonas aerug	inosa op	rC gene fo	r outer m	embrane p	orotein (	C, complete cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10626550_c3_128	986	2906	65	198		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
11017010_c3_139	987	2907	247	744		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12894752_f1_13	988	2908	69	210	84	0.016
Protein name				_	s Name	Acc#
conserved hypothet	ical pro	otein	-	pir:A	72221	A72221
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
13759627_c3_138	989	2909	88	267	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
14633433_f3_55	990	2910	122	369	
Protein name				Locus Name	Acc#
Description					
NO-HIT		·			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
16023402_c3_125	991	2911	887	2664 1410	6.3e-160
Protein name				Locus Name sp:FTSK_COXB	Acc# P39920
Description					
CELL DIVISION PROTE	IN FTS	K HOMOLOG			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19537930_£1_1	992	2912	97	294 116	4.5e-07
Protein name				Locus Name	Acc#
hypothetical protei	n APEOS			pir:D72685	D72685
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
19812500_f1_14	993	2913	974	2925 2125	5.8e-220
Protein name				Locus Name sp:DPO1_HAEI	Acc# N P43741
Description	t				
DNA POLYMERASE I,	(POL I)				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2008433_c1_102	994	2914	287	864		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT				-		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20506502_c2_105	995	2915	62	189		
Protein name				Locu	s Name	Acc#
Description						
по-ніт	······································					
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
20581377_c3_127	996	2916	888	2667	2717	1.1e-282
Protein name				Locu	s <u>Name</u>	Acc#
DNA topoisomerase,				pir:G	64119	G64119
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2131550_f1_9	997	2917	115	348	153	5.4e-11
Protein name					s Name	Acc#
pterin-4-alpha-cark dehydratase:protein			s12296	pir:S	74881	S74881
Description				_1		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22119402_f3_77	998	2918	262	789	491	8.2e-47
Protein name					s Name	Acc# P35115
Description						
OCTOPINE TRANSPORT	SYSTEM	PERMEASE I	PROTEIN O	ССМ	·	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
234408_c2_111	999	2919	129	390	275	6.3e-24
Protein name					s Name MA_ECOLI	Acc# P36999
Description						
METHYLTRANSFERASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score '	Probability
23928130_t3_66	1000	2920	63	192		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24256553_f2_31	1001	2921	751	2256	2080	3.4e-215
Protein name				_	s Name	Acc#
DNA topoisomerase	IV			gp:AB	023570	AB023570
				┙ Ĺ		
<u>Description</u>				J [		
Description Vibrio parahaemoly	/ticus pa	arC gene	for DNA to	poisomera	ase IV,co	omplete cds.
	yticus pa NTID	arC gene	for DNA to  NT  Length	poisomera  AA  Length	se IV,co	omplete cds.  Probability
Vibrio parahaemoly			NT	<u>AA</u>		
Vibrio parahaemoly ORF Name	NTID	AAID	<u>NT</u> Length	AA Length 891	Score	Probability
Vibrio parahaemoly ORF Name 24645763_c1_91	NTID 1002	<u>AAID</u>	<u>NT</u> Length	AA Length 891	Score 255 S Name	Probability 8.4e-22
ORF Name  24645763_c1_91  Protein name	NTID 1002	<u>AAID</u>	<u>NT</u> Length	AA Length 891 Locus	Score 255 S Name	Probability  8.4e-22  Acc#
ORF Name  24645763_c1_91  Protein name hypothetical prote	NTID 1002	<u>AAID</u>	<u>NT</u> Length	AA Length 891 Locus	Score 255 S Name	Probability  8.4e-22  Acc#
ORF Name  24645763_c1_91  Protein name hypothetical protein description	NTID 1002	<u>AAID</u> 2922	NT Length 296	AA Length 891 Locus pir:G	Score 255 s Name 71841	Probability  8.4e-22  Acc# G71841
ORF Name  24645763_c1_91  Protein name hypothetical protein Description  ORF Name	NTID  1002  ein jhp11  NTID	<u>AAID</u> 2922 155 <u>AAID</u>	NT Length 296 NT Length	AA Length  B91  Locus  pir:G  AA Length  213	Score 255 s Name 71841	Probability  8.4e-22  Acc# G71841
ORF Name  24645763_c1_91  Protein name hypothetical protein  Description  ORF Name  24881313_f2_32	NTID  1002  ein jhp11  NTID	<u>AAID</u> 2922 155 <u>AAID</u>	NT Length 296 NT Length	AA Length  B91  Locus  pir:G  AA Length  213	Score 255 S Name 71841 Score	Probability  8.4e-22  Acc# G71841  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
25417950_c3_134	1004	2924	162	489 214	1.8e-17
Protein name				Locus Name sp:RRMA_ECOL	Acc# P36999
Description					
METHYLTRANSFERASE)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
31406308_c2_113	1005	2925	114	345	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
31485625_f3_72	1006	2926	275	828 427	5.0e-40
Protein name				Locus Name	Acc#
				gp:AB032934	AB032934
Description					
Vibrio alginolytic proteins, complete		, orfC, d	orfD genes	for PF60 andhypo	thetical
ORF Name	NTID	AAID.	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
34025462_c1_104	1007	2927	69	210	
Protein name					
Process name				Locus Name	Acc#
<u>Description</u>					Acc#
					Acc#
Description	NTID	AAID	NT Length		Acc# Probability
Description NO-HIT	NTID	AAID 2928		Locus Name  AA Score	
Description NO-HIT ORF Name			Length	Locus Name  AA Length Score	
Description NO-HIT ORF Name 34086012_c1_97			Length	Locus Name  AA Length  417	Probability

ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	AA Length Score	Probability
35360075_f1_2	1009	2929	166	501 73	0.032
Protein name nef protein Description				Locus Name gp:AF169778	Acc# AF169778
HIV-1 isolate G2 long terminal rep			_	ef) gene, partia	1 cds;and 3'
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
36110253_f2_36	1010	2930	108	327 76	0.0077
Protein name  outer surface propertion	otein A	· · · · · · · · · · · · · · · · · · ·	<del></del>	Locus Name	Acc# X68539
	WudII) pla	smid Osp	A gene for	outer surface p	roteinA.
			NT	AA	
ORF Name	NTID	AAID	Length	Length Score	Probability
3945893_f3_71	1011	2931	286	861 825	3.3e-82
Protein name				Locus Name gp:AB032934	Acc# AB032934
Description  Vibrio alginolyt	-	orfC, o	rfD genes	for PF60 andhypo	thetical
proteins, complet	e cas.		NTT	7.7	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
4337963_f2_51	1012	2932	268	807 434	9.0e-41
Protein name				Locus Name gp:AB032934	Acc# AB032934
Description					<del></del>
Vibrio alginolyt proteins, complet	_	orfC, o	rfD genes	for PF60 andhypo	thetical
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4691525_f1_7	1013	2933	78	237	
Protein name				Locus Name	Acc#
Description					
NO-HIT		······································			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4735333_c2_110	1014	2934	570	1713 1749	4.0e-180
Protein name				Locus Name sp:RF3_HAEIN	Acc# . P43928
Description					
PEPTIDE CHAIN RELEA	ASE FACT	OR 3 (RF-	3)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
5102193_t2_54	1015	2935	372	1119 897	7.8e-90
Protein name				Locus Name	Acc# P43889
Description					
ACETYLGLUCOSAMINE-1	-PHOSPI	HATE URIDY	LTRANSFER	ASE)	
ORF Name	NTID	AAID	NT Length	AA Score	Probability
5111013_c3_126	1016	2936	201	606	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
6516518_f1_25	1017	2937	243	732 499	1.2e-47
Protein name				Locus Name sp:NOCQ_AGRT5	Acc# P35118
Description					
NOPALINE TRANSPORT	SYSTEM	PERMEASE	PROTEIN N	OCQ	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
6820875_c1_100	1018	2938	345	1038 159	8.8e-09
Protein name				Locus Name	Acc#
apolipoprotein A-IV	precur	rsor		pir:C40892	C40892
Description					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
807692_c3_137	1019	2939	563	1692	171	1.2e-08
Protein name Trip230					Name 007217	Acc# AF007217
Description						
Homo sapiens Trip	230 mRNA	, complet	e cds.			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
978502_£2_52	1020	2940	266	801	430	2.4e-40
Protein name					S Name 032934	<u>Acc#</u> AB032934
Description						
Vibrio alginolyti proteins, complete	-	, orfC, o	rfD genes	for PF60	andhypot	chetical
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10056500 f1 3	1021	2941	78	237	163	4.7e-12
10030300_11_3		ــــــــــــــا لــ				1
Protein name					s Name	Acc#
					S Name	Acc# B64145
Protein name				Locus	S Name	
Protein name  hypothetical protein  Description  ORF Name	ein HI018	AAID	NT Length	Locus pir:B	Score	B64145 Probability
Protein name hypothetical protein Description	ein HI018	37	NT	Locus pir:B	S Name 64145	B64145
Protein name  hypothetical protein  Description  ORF Name	ein HI018	AAID	NT Length	Locus  AA  Length  1392  Locus	Score  831 S Name	B64145 Probability
Protein name hypothetical protein  ORF Name  10656456_c3_200  Protein name	ein HI018	AAID	NT Length	Locus  AA  Length  1392  Locus	Score  831	Probability 7.7e-83
Protein name hypothetical protein Description ORF Name	ein HI018	AAID	NT Length	Locus  AA  Length  1392  Locus	Score  831 S Name	B64145  Probability 7.7e-83  Acc#
Protein name hypothetical protein  ORF Name  10656456_c3_200  Protein name	NTID	AAID 2942	NT Length 463	Locus  AA  Length  1392  Locus  Sp:YW	SCOTE  831 S Name BN_BACSU	Probability  7.7e-83  Acc# P39597
Protein name hypothetical protein  ORF Name  10656456_c3_200  Protein name  Description	NTID	AAID 2942	NT Length 463	Locus  AA  Length  1392  Locus  Sp:YW	SCOTE  831 S Name BN_BACSU	Probability  7.7e-83  Acc# P39597
Protein name hypothetical protein  ORF Name  10656456_c3_200  Protein name  Description  HYPOTHETICAL 45.7	NTID  1022  KD PROT	AAID 2942 EIN IN EP	NT Length 463	Locus  AA  Length  1392  Locus  Sp:YW	SCORE  831 SNAME BN_BACSU  REGION PR	Probability 7.7e-83  Acc# P39597  RECURSOR
Protein name hypothetical protein  ORF Name  10656456_c3_200  Protein name  Description  HYPOTHETICAL 45.7  ORF Name	NTID  NTID  NTID  NTID  NTID	AAID 2942 EIN IN EP AAID	NT Length 463 R-GALK INT NT Length	Locus  Pir:B  AA  Length  1392  Locus  Sp:YW  ERGENIC I  AA  Length  1677  Locus	SCOTE  BACSU  REGION PR	Probability 7.7e-83 Acc# P39597 RECURSOR Probability
Protein name hypothetical protein  ORF Name  10656456_c3_200  Protein name  Description  HYPOTHETICAL 45.7  ORF Name  12603450_f1_20	NTID  NTID  NTID  NTID  NTID	AAID 2942 EIN IN EP AAID	NT Length 463 R-GALK INT NT Length	Locus  Pir:B  AA  Length  1392  Locus  Sp:YW  ERGENIC I  AA  Length  1677  Locus	SCORE  831 SNAME BN_BACSU  REGION PR  SCORE  1262 S NAME	Probability 7.7e-83  Acc# P39597  RECURSOR  Probability 1.6e-128  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
12978955_£2_43	1024	2944	274	825 609	2.6e-59
Protein name				Locus Name sp:YH25_AZOCH	Acc# P54085
Description					
(ORF5)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
13679786_f3_112	1025	2945	110	333 105	1.2e-05
Protein name				Locus Name	Acc#
hypothetical prote	ein			gp:BSZ75208	Z75208
Description					
B.subtilis genomic	sequen	ce 89009bp	).		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
13712643_c3_187	1026	2946	215	648 113	0.00040
Protein name				Locus Name	Acc#
conserved hypothet	ical pr	otein		pir:B75483	B75483
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
14191915_f2_69	1027	2947	365	1098 329	2.5e-34
Protein name				Locus Name	Acc#
conserved hypothet	ical pr	otein ylbK	•	pir:H69874	H69874
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14251568_f2_72	1028	2948	118	357 207	1.0e-16
Protein name				Locus Name	Acc#
hypothetical prote	ein APE1	486		pir:F72628	F72628
Description				<u> </u>	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14492180_c2_149	1029	2949	67	204		
Protein name				Locus	s_Name	Acc#
Description						
NO-HIT	_					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14511561_c3_197	1030	2950	1319	3960	3473	0.0
Protein name					s Name	Acc#
phosphoribosylformy synthase,:formylglyc synthetase:phosphori	inamide	ribonucl		pir:S	YECPG	D65033:A31 862:A34192
Description			10-10-10-10-10-10-10-10-10-10-10-10-10-1	<b>.</b> _		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14878927_t2_42	1031	2951	271	816	294	4.8e-39
Protein name					s Name	Acc#
<u>Description</u>				sp:HI	S2_AQUAE	O67780
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14882750_c2_164	1032	2952	409	1230	585	9.0e-57
Protein name					s Name	Acc#
putative membrane t	ranspor	t protein	•	gp:SC	C75A	AL133220
Description						
Streptomyces coelic	olor co	smid C75A	•			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14970637_c3_176	1033	2953	237	714	791	1.3e-78
Protein name					s Name	Acc#
Description				ph:CI	PP_ECOLI	P19245
PROTEIN F21.5)		-				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15831636_c2_158	1034	2954	127	384	236	1.5e-19
Protein name Acriflavin resist Description	ance prot	ein D.		Locus gp:D9	s <u>Name</u> 0846	Acc# D90846:AB0 01340
E.coli genomic DN	A, Kohara	clone #3	57 (46.5-4	6.8 min.	٠.	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
165842_c2_157	1035	2955	205	618	332	5.8e-30
Protein name					s Name LH_RHIME	<u>Acc#</u> P25198
Description						
NODULATION PROTEI	N NOLH PR	ECURSOR				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
16687526_f1_10	1036	2956	448	1347	1082	1.9e-109
Protein name  Description				1	s Name GA_ECOLI	P08205:068 009:068010 :068011:06
SYNTHASE) (AGS)						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
191675_c2_150	1037	2957	251	756	698	9.5e-69
Protein name 5 adenylylsulfat	e APS red	uctase		_	Name 170343	Acc# AF170343
Description						
Burkholderia cepa cds; and ATP sulfu						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
19821942_c3_205	1038	2958	64	195		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20203302_c3_185	1039	2959	226	681 141	4.0e-08
DnrE protein Description			, <u>-</u> -	Locus Name gp:PST131716	Acc# AJ131716
Pseudomonas stutze	eri dnrE	gene and	ORF235 (p	artial).	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20570385_c2_159	1040	2960	193	582	
Protein name				Locus Name	Acc#
Description					
NO-HIT		<del></del>			
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
21494010_c2_174	1041	2961	455	1368 1377	1.1e-140
Protein name				Locus Name	Acc#
nitric oxide reduc	tase			gp:AF002217	AF002217
Description					
Ralstonia eutropha	a megapla	asmid pHG1	nitric c	xide reductase	(norB)gene,
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
21676712_c1_120	1042	2962	130	393 395	1.2e-36
Protein name				Locus Name	Acc#
sulfate adenylylti	ransferas	se subunit	: CysN	gp:AF130466	AF130466
Description					
Campylobacter jeji alpha-2,3-sialyltra CysD (cysD) genes, (cysN) gene, partia	ansferas complet	e (cst-I)	and sulfa	teadenylyltransf	erase subunit

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22078812_c2_154	1043	2963	727	2184	1636	3.8e-168
Protein name  Description					s Name CG_ECOLI	Acc# P24230:P76 721
ATP-DEPENDENT DNA F	ELICAS	E RECG,				
ORF Name 22350926 cl 115	NTID	AAID 2964	NT Length	AA Length 2175	Score	Probability 3.6e-227
Protein name		] [	J []	Locu	s Name OB_PSEFR	Acc# P28793
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
23556252_c3_203	1045	2965	600	1803	709	6.0e-85
Protein name	/	(marie - )		_	s Name	Acc#
glutamate synthase Description	(Terrec		molog yeld	pir:C	69794	C69794
ORF Name 23593830_c3_199	NTID 1046	<u>AAID</u>	NT Length	AA Length 1023	Score	Probability 9.7e-44
Protein name					s Name BM_BACSU	Acc# P39596
Description				<u></u>		
HYPOTHETICAL 42.8 F	D PROTE	EIN IN EPI	R-GALK INT	'ERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23651900_f2_48	1047	2967	328	987	634	5.8e-62
Protein name					s Name HI_HAEIN	Acc# P44606
Description						
HYPOTHETICAL PROTE	N HI02	70			-	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabil	lity
23862576_c1_124	1048	2968	153	462	182	4.5e-14	
Protein name probable antibiotic	regist	ance prof	ein mtrC	_	Name		Acc#
Description		ance proc		pir:S		S4:	2418:S40 2
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probabil	ity
24042500_c3_183	1049	2969	313	942	608	3.3e-59	
Protein name					Name N_MYCTU		<u>lcc#</u> 10600
Description							
SULFURYLASE)							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabil	.ity
24302260_c3_179	1050	2970	410	1233	1331	8.0e-136	
Protein name 3-oxoacyl-CoA thio	lase			_	Name 150672		Acc# F150672
<del></del>	lase			_			
3-oxoacyl-CoA thio		cyl-CoA th	iolase (f	gp:AF	150672	Al	
3-oxoacyl-CoA thio		cyl-CoA th	iolase (f <u>NT</u> Length	gp:AF	150672	Al	F150672
3-oxoacyl-CoA thiol Description Pseudomonas putida	3-охоас		NT	gp:AF	150672 e, comple	Aletecds.	F150672
3-oxoacyl-CoA thiol Description Pseudomonas putida ORF Name	3-oxoac	AAID	<u>NT</u> Length	gp:AF:  adA) gene  AA  Length  432  Locus	150672 e, comple	Probabil	F150672
3-oxoacyl-CoA thiol Description Pseudomonas putida ORF Name 25584438_c3_189 Protein name	3-oxoac	AAID	<u>NT</u> Length	gp:AF:  adA) gene  AA  Length  432  Locus	s Name	Probabil	ity
3-oxoacyl-CoA thiol Description Pseudomonas putida  ORF Name 25584438_c3_189 Protein name CeoB	3-0x0ac <u>NTID</u> 1051	AAID 2971	NT Length 143	gp:AF:  adA) gene  AA  Length  432  Locus  gp:BC	s Name	Probabil 7.6e-12	ity
3-oxoacyl-CoA thiol Description Pseudomonas putida  ORF Name 25584438_c3_189 Protein name CeoB Description	3-0x0ac <u>NTID</u> 1051	AAID 2971	NT Length 143	gp:AF:  adA) gene  AA  Length  432  Locus  gp:BC	s Name	Probabil 7.6e-12	ity Acc#
3-oxoacyl-CoA thiol Description Pseudomonas putida  ORF Name 25584438_c3_189 Protein name CeoB Description Burkholderia cepac	3-0x0ac  NTID  1051	AAID 2971 (ceoA) and	NT Length 143 d CeoB (c	adA) gene  AA  Length  432  Locus  gp:BCC  eoB) gene	s Name J97042 es, comple	Probabil 7.6e-12  Us	ity Acc#
3-oxoacyl-CoA thiol Description Pseudomonas putida  ORF Name 25584438_c3_189 Protein name CeoB Description Burkholderia cepac	3-0x0ac  NTID  1051  ia CeoA  NTID  1052	AAID (ceoA) and AAID 2972	NT Length  143  d CeoB (c  NT Length  134	adA) general AA Length Gp:BCC GoB) general AA Length AA Length AD Locus	Score  174 S Name 197042 Score  133 S Name	Probabil  7.6e-12  Usetecds.  Probabil  1.8e-07	ity Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	ability
26741556_c1_135	1053	2973	198	597	101	0.001	.1
Protein name		,			s Name 34_BRELC		<u>Acc#</u> Q99074
Description							
HAM34 PROTEIN							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
3245640_c3_190	1054	2974	425	1278	265	9.4e-	20
Protein name				Locu	s Name		Acc#
probable cation eff	lux sys	stem prote	ein	pir:E	71874		E71874
Description				<u></u>			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
3375052_c1_125	1055	2975	144	435	177	3.6e-	12
Protein name				Locu	s_Name		Acc#
probable efflux tra	insporte	er		pir:H	71918		H71918
Description				<u> </u>			
ORF Name	NTID	AAID	NT Length	AA Length	Score		bility
34027092_c2_175	1056	2976	74	222	60	0.025	
Protein name				Locu	s Name		Acc#
tonoplast intrinsic	: protei	n		gp:AF	037061		AF037061
Description				<del></del>			
Zea mays tonoplast	intrins	sic protei	in (ZmTIP1	) mRNA,	complete	cds.	·
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
35197126_f3_79	1057	2977	179	540	155	3.3e-	11
Protein name				Locus	s Name		Acc#
TatB protein				gp:EC	05830		AJ005830
Description					,		
Escherichia coli ta	atABCD o	operon.					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35339135_t2_61	1058	2978	264	795	308	2.9e-35
Protein name				-	s Name P3_AERHY	Acc# P45794
Description		-				
TYPE 4 PREPILIN-LI	KE PROTI	EIN SPECIF	IC LEADER	PEPTIDAS	SE,	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35395926_f1_6	1059	2979	511	1536	325	2.2e-35
Protein name				Locus	Name	Acc#
probable helicase				pir:T	40239	T40239
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36047308_£2_50	1060	2980	105	318		
Protein name				Locus	<u>Name</u>	Acc#
Description						
NO-HIT		<del></del>				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3939838_c2_173	1061	2981	66	201	189	8.2e-15
Protein name					Name 35_PSESY	Acc# P52830
Description						
50S RIBOSOMAL PROT	EIN L35					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3947675_f2_46	1062	2982	250	753	375	1.6e-34
Protein name					Name	Acc# P12380
Description						
HISTIDINE UTILIZAT	ION REP	RESSOR				

ORF Name	NTID	AAID	<u>NT</u> Length	Length Score	Probability		
3954218_c3_177	1063	2983	441	1326 1252	7.6e-140		
Protein name				Locus Name	Acc# P44838		
Description							
ATP-DEPENDENT CL	P PROTEAS	E ATP-BIN	DING SUBUN	IT CLPX			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability		
4328428_c1_119	1064	2984	310	933 911	2.6e-91		
Protein name				Locus Name sp:CYSD_MYCTU	Acc# Q10599		
Description							
SULFURYLASE)							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability		
4485963_c3_206	1065	2985	121	366 469	1.8e-44		
Protein name				Locus Name	Acc#		
ribosomal protein	n L20 ———			pir:R5EC20	D64930:S08		
<u>Description</u>					608:A02806 :I41282		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability		
5120443_f3_94	1066	2986	214	645 363	3.0e-33		
Protein name				Locus Name sp:YACE_VIBVU	Acc# Q56741		
Description							
HYPOTHETICAL 22.5 KD PROTEIN IN VVPD 3'REGION (ORFX)							
	5 KD PROT	EIN IN VV	PD 3'REGIO	N (ORFX)			
ORF Name	5 KD PROT	AAID	PD 3'REGIO  NT  Length	N (ORFX)  AA  Length  Score	Probability		
ORF Name 5890643_t2_73			NT	AA Score	Probability 7.7e-07		
	NTID 1067	AAID	<u>NT</u> Length	AA Score			

ORF Name	NTID Z	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
7277_£3_93	1068	2988	423	1272	664	3.8e-6	55
Protein name pilus assembly pr	otein PilC			_	3 Name 038655		<u>Acc#</u> AF038655
Description				_			
Legionella pneumo protein PilC (pilC peptidase PilD (pi	), and type	e IV prep	oilin-lik				-
ORF Name	NTID A	AID	<u>NT</u> Length	AA Length	Score	Probal	oility
978517_c2_166	1069	2989	433	1302	615	5.8e-1	112
Protein name				-	s <u>Name</u> TS_HAEIN		Acc# P45240
Description							
SODIUM/GLUTAMATE	SYMPORT CAP	RRIER PRO	TEIN (GL	UTAMATE I	PERMEASE)		<u></u>
ORF Name	NTID A	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
10547711_c2_8	1070	2990	275	828	933	1.2e-9	93
Protein name					Name C_HAEIN		<u>Acc#</u> P44785
Description						-	
ATP-BINDING PROTE	IN ABC						
ORF Name		AAID	NT Length	AA Length	Score	Probab	
188807_c2_10	1071	2991	110	354	320	1.16-2	.0
Protein name					Name		Acc#
Description				SD: 511	PA_PASHA 		Q08868:Q07 363
OUTER MEMBRANE L	POPROTEIN 1	PRECURS	OR (PLP1	)			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19589635_c1_7	1072	2992	99	300 190	6.5e-15
Protein name ORF120				Locus Name	Acc# D15061
Description					
E.coli genomic DN	A, 5'fla	nking reg	gion of rrr	M gene.	
ORF Name	NTID	AAID	<u>NT</u> Length	$\frac{\underline{AA}}{\underline{Length}}  \frac{\underline{Score}}{}$	Probability
24353193_f1_1	1073	2993	95	288 113	9.3e-07
Protein name				Locus Name	Acc#
hypothetical prote	ein PH01	33		pir:C71234	` C71234
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
35582056_c3_11	1074	2994	240	723 631	1.2e-61
Protein name				Locus Name sp:YAEE_HAEIN	Acc# P46492
Description				•	
HYPOTHETICAL ABC	TRANSPOR	TER PERMI	EASE PROTEI	N HI0620.1	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4144442_f2_6	1075	2995	131	396	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16924127_f1_1	1076	2996	367	1104 137	7.2e-06
Protein name				Locus Name	<u>Acc#</u> M60717
Description				L	
P.stutzeri NosA p	rotein (	nosA) ger	ne, complet	e cds.	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
5109792_f2_3	1077	2997	122	369	
Protein name				Locus Name	Acc#
Description					
NO-HIT	-				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16681688_c1_3	1078	2998	60	183 55	0.044
Protein name				Locus Name sp:RNH_HELPY	Acc# P56120
Description					
RIBONUCLEASE H, (	RNASE H)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24039143_f1_1	1079	2999	345	1038 637	2.8e-62
Protein name				Locus Name	Acc#
ornithine decarbo	xylase			pir:D72200	D72200
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29337840_f2_2	1080	3000	150	450 95	0.014
Protein name				Locus Name	Acc#
AvtA		- '-		gp:AF014804	AF014804
Description					
Neisseria meningi genes, complete cd		IB (pglB),	PgIC (pg	IC), PglD (pglD	), andAvtA (avtA)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
11900461_f2_7	1081	3001	69	210	
Protein name				Locus Name	Acc#
Description					
NO-HIT					

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
14573562_±1_2	1082	3002	341	1026	1.1e-108
Protein name				Locus Name	
<u>Description</u>					P25745:P75 964
(EC 2.1.1.61)					
ORF Name	NTID	AAID	NT Length	Length Score	Probability
30270465_f1_3	1083	3003	249	750 185	5.7e-14
Protein name				Locus Name sp:YYAD_BACSU	Acc# P37520
Description					
HYPOTHETICAL 37.	7 KD PROT	EIN IN RP	SF-SPOOJ I	NTERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
6365936_£2_6	1084	3004	108	327 170	8.5e-13
					<b>-</b>
Protein name		<b>-</b>		Locus Name gp:ECPURB	Acc# X59307
Protein name  Description					
	RF-23, pu	rB and ph	oP (5'end)	gp:ECPURB	
Description  E.coli ORF-15, O	RF-23, pu:	rB and ph	<u>NT</u> Length	gp:ECPURB	X59307  Probability
Description E.coli ORF-15, O			NT	gp:ECPURB  genes.  AA Score	X59307
Description  E.coli ORF-15, O	NTID	AAID	<u>NT</u> Length	gp:ECPURB  genes.  AA Length Score	X59307  Probability
Description  E.coli ORF-15, O  ORF Name  6537957_c1_11	NTID	AAID	<u>NT</u> Length	genes.  AA Score Length  603  454  Locus Name	Probability  6.8e-43  Acc#
Description  E.coli ORF-15, O  ORF Name  6537957_c1_11  Protein name	NTID 1085	AAID 3005	NT Length 200	gp:ECPURB  genes.  AA Score Length 603 454  Locus Name sp:YGBB_ECOLI	Probability  6.8e-43  Acc# P36663
Description  E.coli ORF-15, O  ORF Name  6537957_c1_11  Protein name  Description	NTID 1085	AAID 3005 EIN IN SU AAID	NT Length 200	genes.  AA Score Length 454  Locus Name Sp:YGBB_ECOLI  TERGENIC REGION  AA Score	Probability  6.8e-43  Acc# P36663
Description  E.coli ORF-15, O.  ORF Name  6537957_c1_11  Protein name  Description  HYPOTHETICAL 16.	NTID 1085	AAID 3005 EIN IN SU	NT Length 200 RE-CYSC IN	genes.  AA Score  Length  603 454  Locus Name  sp:YGBB_ECOLI  TERGENIC REGION  AA Score	Probability  6.8e-43  Acc# P36663
Description  E.coli ORF-15, O.  ORF Name  6537957_c1_11  Protein name  Description  HYPOTHETICAL 16.  ORF Name	NTID  1085  9 KD PROT	AAID 3005 EIN IN SU AAID	NT Length 200 RE-CYSC IN NT Length	genes.  AA Score Length 454  Locus Name Sp:YGBB_ECOLI  TERGENIC REGION  AA Score	Probability  6.8e-43  Acc# P36663
Description  E.coli ORF-15, O.  ORF Name  6537957_c1_11  Protein name  Description  HYPOTHETICAL 16.  ORF Name  173187_f3_5	NTID  1085  9 KD PROT	AAID 3005 EIN IN SU AAID	NT Length 200 RE-CYSC IN NT Length	genes.  AA Score Length 454  Locus Name Sp:YGBB_ECOLI  TERGENIC REGION  AA Score Length Score	Probability  6.8e-43  Acc# P36663  (ORF0)  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34407827_f2_2	1087	3007	366	1101	786	4.5e-78
Protein name					s Name FA_ECOLI	Acc# P29212
Description						
SYNTHETASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
7285152_f1_1	1088	3008	69	207		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16056563_c2_63	1089	3009	141	426	90	0.020
Protein name					s Name	Acc#
hypothetical wtfw p	protein			pir:T	41252	T41252
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1986087_f3_30	1090	3010	412	1239	1316	3.1e-134
Protein name					s Name RA_HAEIN	Acc# P43885
Description				L		
D-3-PHOSPHOGLYCERA	TE DEHY	DROGENASE,	(PGDH)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2072762_f1_10	1091	3011	1220	3663	524	4.4e-93
Protein name					s Name	Acc#
chromosome segregat	tion SM	C		pir:G	69708	
DIOCEIN: MINICHTOMOS			rotein	<u>                                   </u>		
SMC			rotein			G69708:JC4 819:PC4029

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21531252_f3_36	1092	3012	303	912	722	2.7e-71
Protein name				Locus	s Name	Acc#
translation elong	ation fac	ctor EF-Ts		pir:E	FECS	
Description					_	A03525:A45 269:A32881
						:S45235:B6
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
23406_f1_11	1093	3013	278	837	898	6.1e-90
Protein name				Locus	s Name	Acc#
				sp:RS	2_SPIPL	P34831
<u>Description</u>				-		
30S RIBOSOMAL PRO	TEIN S2					
			NT	AA		December 12 days
ORF Name	NTID	AAID	<u>Length</u>	Length	Score	Probability
23437562_f1_7	1094	3014	253	762	186	3.2e-18
Protein name				Locus	s Name	Acc#
hypothetical prot	ein HP086	52		pir:F	64627	F64627
<u>Description</u>						······································
			NT	AA		
ORF Name	NTID	AAID	Length	Length	Score	Probability
23613510_c2_70	1095	3015	374	1125	700	5.8e-69
Protein name				Locus	s Name	Acc#
				sp:YC	FO_ECOLI	P75949
Description						
HYPOTHETICAL 37.6	KD PROTE	EIN IN FHU	E-NDH INT	ERGENIC I	REGION	
ORF Name	NTID	AAID	NT	AA	Score	Probability
			Length	Length		
23861686_f3_35	1096	3016	180	543	526	1.6e-50
Protein name					Name	Acc#
invasion protein	nomorog			gp:AF	116285	AF116285
Description						
Pseudomonas aerug						
andphosphoenolpyrucds.	vate-prot	ein phosp	notransfe	rase PtsI	(ptsP)	genes, complete

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2460181_f1_1	1097	3017	987	2964	3662	0.0
Protein name					s <u>Name</u> OB_PSEPU	Acc# P19175
Description						
BETA CHAIN) (RNA	POLYMERA	SE BETA SU	BUNIT)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25401687_c3_88	1098	3018	141	426	372	3.3e-34
Protein name					s <u>Name</u> U89892	Acc# U89892
Description				<u></u>		
Pseudomonas aerug	ginosa vi:	rulence fa	ctor regu	lator (v:	fr) gene,	partial cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26064760_c3_93	1099	3019	66	201		
<u>Protein name</u>				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29890701_f3_31	1100	3020	469	1410	1678	1.3e-172
Protein name				Locus	s Name	Acc#
				sp:GS	HR_HAEIN	P43783
Description						
GLUTATHIONE REDUC	CTASE, (GI	R) (GRASE)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
32287567_f2_16	1101	3021	1422	4269	4932	0.0
Protein name  99% identity over coli	1407 ami	no acids	with E.		S Name YSTMF1	Acc# AF170176
Description			- 1	_		
Salmonella typhir	nurium fra	agment STM	F1.		•	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3907166_£1_6	1102	3022	309	930	117	1.6e-06
Protein name  putative biotin pro	otein li	lgase			s <u>Name</u> 016461	Acc# AF016461
Description						<del></del>
Bordetella pertuss and Bvg accessory fa	-		-	_	(birA) ge	ene,complete cds
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4507138_f1_5	1103	3023	727	2184	736	2.5e-88
Protein name					s Name C_ECOLI	Acc# P23865
Description						
PROTEIN)						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
4959501_c1_46	1104	3024	83	252		
Protein name	•			Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
5119452_c3_92	1105	3025	267	804	398	5.9e-37
Protein name					s <u>Name</u> 02_HAEIN	Acc# P44070
Description						
HYPOTHETICAL PROTE	IN HI090	02				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
11955251_c2_26	1106	3026	915	2748	2735	1.3e-284
Protein name					S Name A ECOLI	Acc#
Description						P00957:P78
ALANYL-TRNA SYNTHE	rase, (7	ALANINET	RNA LIGAS	E) (ALARS	5)	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
16523292_c2_25	1107	3027	488	1467	1534	2.4e-	157
Protein name					s Name R8_HAEIN		<u>Acc#</u> P44797
Description							
ADENYLOSUCCINATE LY	ASE, (A	DENYLOSUC	CINASE) (	ASL)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
23475417_£3_18	1108	3028	73	222			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT			<del> </del>		<del></del>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
24308500_c1_22	1109	3029	155	468	282	1.2e-	24
<u>Protein name</u>				Locus	Name		Acc#
erythroid different	iation-	related fa	actor 2	gp:AF	040248		AF040248
Description							
Homo sapiens erythr	oid dif	ferentiat:	ion-relat	ed factor	2 mRNA,	partia	ıl cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
31444625_c3_34	1110	3030	452	1359	646	3.1e-	63
<u>Protein name</u>					Name LF_BACSU		<u>Acc#</u> P94408
Description							
HYPOTHETICAL 53.3 K	D PROTE	IN IN SFP	-GERKA IN	TERGENIC	REGION		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
33632752_c2_27	1111	3031	285	855	679	9.8e-	67
Protein name aspartate kinase, I precursor:lysine-sen		aspartokir	nase II	Locus pir:A	Name 18946		Acc# A48946:B48
Description							946:C48946

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
33829043_c3_33	1112	3032	106	321	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4400283_c1_23	1113	3033	95	288	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
6852262_c1_21	1114	3034	61	186	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
9933552_£1_3	1115	3035	82	249	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
9975052_c2_24	1116	3036	132	399	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
1036637_c3_276	1117	3037	61	186	
Protein name				Locus Name	Acc#
Description					
NO-HIT				-	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1038885_f1_26	1118	3038	416	1251	1526	1.7e-156
Protein name lactate dehydrogena	se			_	<u>Name</u> U58911	Acc# U58911
Description						
Neisseria meningiti complete cds, HI1054					HI0379 H	nomologgenes,
			NT	AA		
ORF Name	NTID	AAID	Length	Length	Score	Probability
10547558_c1_180	1119	3039	176	531	123	8.1e-08
Protein name				Locus	s Name	Acc#
hypothetical protei	n APE11	65		pir:H	72586	H72586
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
11711_c3_265	1120	3040	434	1305	489	1.3e-46
Protein name				Locus	s Name	Acc#
HisX				gp:AF	010189	AF010189
Description		-				
Pseudomonas stutzer complete cds; and Pu		_	_		HisX (his	sX)gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
11991552_f3_108	1121	3041	286	861	614	7.6e-60
Protein name				Locus	s Name	Acc#
				sp:TR	PC_PSEPU	P20578
Description						
INDOLE-3-GLYCEROL P	ноѕрнат	E SYNTHAS	SE, (IGPS)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12222077_f2_53	1122	3042	198	597	469	1.8e-44
Protein name				Locus	Name	Acc#
•				sp:RE	CR_HAEIN	P44712
Description						
RECOMBINATION PROTE	IN RECR					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
136302_c1_167	1123	3043	154	465	95	0.0012
Protein name hypothetical protei	n sll16	75		Locus pir:S	8 Name 74649	<u>Acc#</u> S74649
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
13678763_c1_182	1124	3044	223	672	194	2.4e-15
Protein name hypothetical protei	n RP471	-		Locus pir:D	<u> Name</u> 71706	<u>Acc#</u> D71706
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14094452_c3_267	1125	3045	60	183		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14103402_f1_7	1126	3046	286	861	265	7.3e-23
Protein name					s Name UG_BACSU	Acc# P35154
Description						
HYPOTHETICAL 29.6 F	D PROTI	IN IN RIB	T-DACB IN	TERGENIC	REGION	(ORFX7)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14273586_t2_57	1127	3047	451	1356	1508	1.4e-154
<u>Protein name</u>					s Name CC_PSEAE	Acc# P37798
Description						<b>_</b>
CARBOXYLASE,) (ACC)						·

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14572132_f3_135	1128	3048	238	717	736	8.9e-73
Protein name  Description				-	Name D3_HAEIN	Acc# P44319
LYASE)						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
14875305_±3_113	1129	3049	139	420	141	1.2e-09
Protein name Ribonuclease D (	EC 3.1.13.	<del>-)</del>		Locus gp:D90	Name 0825	Acc#
Description						D90825:AB0 01340
E.coli genomic D	NA, Kohara	clone #3	34 (40.6-4	1.0 min.)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16507827_c3_256	1130	3050	519	1560	1030	6.3e-104
Protein name					Name	Acc#
Description				sp:NAI	OB_PSEAE	Q51363:Q51 412
L-ASPARTATE OXID	ASE, (QUIN	OLINATE S	SYNTHETASE	B)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16510254_c1_154	1131	3051	308	927	1063	2.0e-107
Protein name	·				Name L ECOLI	Acc#
Description				Sp.Rr.		P07011:P77 340
PEPTIDE CHAIN RE	LEASE FACT	OR 1 (RF-	-1)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16829177_c1_172	1132	3052	95	288	83	0.041
Protein name			· ·	_	Name	Acc#
Description				] Bb:wc	002130	AC002130
The sequence of sequence.	BAC F1N21	from Arab	oidopsis t	haliana d	hromosom	e 1,complete

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19562800_£2_52	1133	3053	120	363	273	1.0e-23
Protein name				-	s <u>Name</u> AB_HAEIN	Acc# P44711
Description				-		
HYPOTHETICAL PRO	TEIN HI044	2				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20312551_f1_8	1134	3054	202	609	243	1.6e-20
Protein name					S Name UH_BACSU	Acc# P35155
Description						
HYPOTHETICAL 22.	0 KD PROTE	IN IN RIB	r-dacb in	TERGENIC	REGION	(ORFX8)
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
20573252_c2_229	1135	3055	205	618	309	1.6e-27
Protein name					Name JA_ECOLI	Acc# P24250
Description						
HYPOTHETICAL 20.	1 KD PROTE	IN IN SEL	O-SPPA IN	TERGENIC	REGION	(ORF183)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21667027_f2_89	1136	3056	832	2499	2093	1.4e-216
Protein name					Name N_ERWAM	Acc# P46067
Description						
ATP-DEPENDENT PRO	OTEASE LA,					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21681503_f1_9	1137	3057	329	990	758	4.2e-75
Protein name					Name	Acc# P45104
Description						I
HYPOTHETICAL PRO	TEIN HI119	9				7

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21754681_f3_119	1138	3058	197	594	284	7.1e-25
Protein name					s Name EY_ECOLI	<u>Acc#</u> P77385
Description						
HYPOTHETICAL 17.5	KD PROT	EIN IN CU	TE-ASNB IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22128380_f1_41	1139	3059	87	264		
Protein name Description				Locus	s Name	Acc#
NO-HIT						]
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
22384635_f1_31	1140	3060	190	573	604	8.7e-59
Protein name HemO		- Constitution			s Name 133695	Acc# AF133695
Description						<u>.</u>
Neisseria meningit partial cds.	idis He	mO (hemO)	gene, com	plete cd:	s; and Hm	nbR(hmbR) gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23444377_c1_199	1141	3061	492	1479	2364	2.7e-245
Protein name				_	s Name	Acc#
outer membrane pro	tein E			gp:MB0	OOMPE	L31788
Description						
Moraxella catarrha	lis out	er membra	ne protein	E gene,	complete	cds.
ORF Name	NTID	<u>AAID</u>	NT Length	AA Length	Score	Probability
234700_c3_268	1142	3062	344	1035	1002	5.8e-101
Protein name				_	s Name	Acc#
unknown				gp:AF	109131	AF109131
Description						
Sinorhizobium meli andmaleylacetoaceta						dunknown gene.

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
23620263_c3_271	1143	3063	267	804 534	2.3e-51
Protein name				Locus Name sp:KDSB_ECOLI	Acc# P04951
Description					
SYNTHETASE) (CMP-	2-KETO-3	-DEOXYOCTO	DLOSONIC A	CID SYNTHETASE)	(CKS)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23651513_f1_43	1144	3064	259	780 589	3.4e-57
Protein name				Locus Name sp:NADC_RHORU	Acc# P77938
Description					
) (QAPRTASE)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24031586_£1_5	1145	3065	383	1152 976	3.3e-98
Protein name				Locus Name sp:TRPD_ACICA	Acc# P00500
Description					
ANTHRANILATE PHOS	PHORIBOS	YLTRANSFER	RASE,		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24397555_f3_140	1146	3066	219	660 165	1.2e-11
Protein name				Locus Name	Acc#
probable corA pro	tein			pir:F70952	F70952
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24415782_f3_106	1147	3067	61	186 87	0.0079
Protein name				Locus Name	Acc#
UUP protein				gp:ECUUP	Y09439
Description					
E.coli uup gene,	partial.			•	

ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
24721962_c3_282	1148	3068	77	234		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24801937_c2_236	1149	3069	63	192		
Protein name				Locus	Name	Acc#
Description						
NO-HIT	· · · · ·					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26756325_f1_1	1150	3070	314	945	1021	5.6e-103
Protein name					Name CA_PSESH	Acc# Q02047
Description	,					
(EC 2.1.3.3) (OT	CASE)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26757790_c3_255	1151	3071	376	1131	1065	1.2e-107
Protein name					Name DA ECOLI	Acc#
Description				25		P11458:P77
QUINOLINATE SYNTE	HETASE A					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
2742036_£1_3	1152	3072	144	435	426	6.3e-40
Protein name					Name ID BACSU	Acc# P52999
Description					_	
DECARBOXYLASE)		<u>,</u>				

ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	AA Length Score	Probability
29301691_£1_14	1153	3073	248	747 140	4.7e-07
Protein name				Locus Name sp:RND_HAEIN	Acc# P44442
Description					
RIBONUCLEASE D, (R	NASE D)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29355287_£2_99	1154	3074	74	225 79	0.013
Protein name				Locus Name	Acc#
MutT/nudix family	protein			pir:A75550	A75550
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29484626_c1_173	1155	3075	356	1071 555	2.4e-57
Protein name				Locus Name	Acc#
				sp:YGI2_PSEPU	P31857
Description					
HYPOTHETICAL 32.4	KD PROT	EIN IN GIL	B-UNCI IN	TERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29531392_f1_40	1156	3076	124	375 173	1.5e-11
Protein name				Locus Name	Acc#
lustrin A				pir:T08852	T08852
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29572155_f2_49	1157	3077	343	1032 406	8.3e-38
Protein name				Locus Name	Acc#
Description				sp:HTRB_HAEIN	P45239:Q48 045
PROTEIN B)				<u></u>	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
29890631_f2_56	1158	3078	157	474 259	3.1e-22
Protein name				Locus Name sp:BCCP_HAEIN	Acc# P43874
Description					<del></del>
BIOTIN CARBOXYL CAR	RIER P	ROTEIN OF	ACETYL-CO	A CARBOXYLASE (B	CCP)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
30204837_c2_230	1159	3079	447	1344 628	2.5e-61
Protein name				Locus Name sp:GPDA_ECOLI	Acc# P37606
Description					
,					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
30330056_£2_48	1160	3080	233	702 506	2.1e-48
Protein name				Locus Name	Acc#
putative ATP-bindin	g prote	ein 		gp:NME242841	AJ242841
putative ATP-bindin	g prote	ein 		gp:NME242841	AJ242841
			region,		AJ242841
Description			region, <u>NT</u> <u>Length</u>		Probability
Description Neisseria meningiti	dis DN	A for opcA	NT	strain Z2491.	
Description Neisseria meningiti ORF Name	dis DNA	A for opcA	NT Length	strain Z2491.  AA Score Length 89  Locus Name	Probability  0.00033  Acc#
Description Neisseria meningiti ORF Name 32878_c2_250	dis DNA	A for opcA	NT Length	strain Z2491.  AA Length Score 243 89	Probability  0.00033  Acc#
Description  Neisseria meningiti  ORF Name  32878_c2_250  Protein name	dis DNA	A for opcA	NT Length	strain Z2491.  AA Score Length 89  Locus Name	Probability  0.00033  Acc#
Description  Neisseria meningiti  ORF Name  32878_c2_250  Protein name  Description	dis DNA	A for opcA	NT Length	strain Z2491.  AA Score Length 89  Locus Name	Probability  0.00033  Acc#
Description  Neisseria meningiti  ORF Name  32878_c2_250  Protein name  Description  SLYX PROTEIN	MTID	A for opcA  AAID  3081	NT Length 80	strain Z2491.  AA Score  Length 89  Locus Name  Sp:SLYX_ECOLI	Probability  0.00033  Acc# P30857
Description  Neisseria meningiti  ORF Name  32878_c2_250  Protein name  Description  SLYX PROTEIN  ORF Name	MTID	A for opcA  AAID  AAID  AAID	NT Length 80 NT Length	strain Z2491.  AA Score Length 89  Locus Name Sp:SLYX_ECOLI  AA Length Score	Probability  0.00033  Acc# P30857  Probability  2.7e-39  Acc#
Description  Neisseria meningiti  ORF Name  32878_c2_250  Protein name  Description  SLYX PROTEIN  ORF Name  34180317_f1_22	MTID	A for opcA  AAID  AAID  AAID	NT Length 80 NT Length	strain Z2491.  AA Score  Length 89  Locus Name  Sp:SLYX_ECOLI  AA Score  Length 420  Locus Name	Probability  0.00033  Acc# P30857  Probability  2.7e-39  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
34417090_f3_151	1163	3083	76	231	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
35196902_f3_129	1164	3084	616	1851 631	1.2e-61
Protein name L-lactate permease	(lctP)	homolog		Locus Name	Acc# F69350
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
35335137_c1_175	1165	3085	330	993 302	8.7e-27
Protein name				Locus Name	·
Description					
DNA POLYMERASE III,	DELTA	SUBUNIT,			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
35353443_c2_215	1166	3086	347	1044 1128	2.6e-114
Protein name				Locus Name	Acc#
				sp:PURA_VIE	PA P40607
Description				sp:PURA_VIE	P40607
Description ADENYLOSUCCINATE SY	NTHETAS	SE, (IMP)	ASPARTATE		PA P40607
ADENYLOSUCCINATE SY	NTID	AAID	<u>NT</u> Length	LIGASE)  AA Length Score	
ADENYLOSUCCINATE SY		-	NT	LIGASE)  AA Score	
ADENYLOSUCCINATE SY	NTID	AAID	<u>NT</u> Length	LIGASE)  AA Length Score	Probability
ADENYLOSUCCINATE SY ORF Name 35603128_f3_150	NTID	AAID	<u>NT</u> Length	LIGASE)  AA Length  243	Probability

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3914143_c2_209	1168	3088	185	558	256	2.1e-29
Protein name ExbB protein				Locus gp:BPE		Acc# AJ132741
Description				-		
Bordetella pertus: ORF1(partial).	sis hupB	, tonB, ex	kbB, exbD	and basR o	genes ar	nd
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3914811_c3_280	1169	3089	370	1113	873	2.7e-87
Protein name				Locus sp:YHCM	Name M_ECOLI	Acc# P46442
Description						
HYPOTHETICAL 43.1	KD PROT	EIN IN RPI	M-HHOA IN	TERGENIC F	REGION	(F375)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3937518_c1_171	1170	3090	264	795	618	2.9e-60
Protein name				Locus	Name	Acc#
Description HYPOTHETICAL 28.9	אר אר אר	CTN TN CT	DE TINGT TA		L_PSEPU	P31856
HYPOTHETICAL 28.9	KD PROT	EIN IN GII			CEGION -	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3938818_c1_166	1171	3091	65	198	228	6.1e-19
Protein name				Locus	Name	Acc#
PurA	_			gp:AF01	L0189	AF010189
Description						
Pseudomonas stutze complete cds; and		_			sX (his	SX) gene,
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3946931_f3_114	1172	3092	216	651	560	4.0e-54
Protein name				Locus gp:ECU8		<u>Acc#</u> U89166
Description						
Eikenella corrode	ns lysin	e decarbo	cylase (EC	ORLD) gene	e, compl	etecds.

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
3960260_f1_23	1173	3093	253	762	282	1.2e-24
Protein name				Locu	s Name	Acc#
probable transcript	ion reg	gulator		pir:T	34763	T34763
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4022217_f2_68	1174	3094	160	483	86	0.00087
Protein name				_	s Name	Acc#
hypothetical protei	n F53A9	9.8		pir:T	16439	T16439
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4023443_c3_278	1175	3095	146	441	498	1.5e-47
Protein name					s Name	Acc#
Description NUCLEOSIDE DIPHOSPH	IATE KII	NASE, (NDK	) (NDP KI		K_PSEAE	Q59636
			NT	AA		
ORF Name	NTID	AAID	<u>Length</u>	<u>Length</u>	Score	Probability
4101387_f1_4	1176	3096	219	660	710	5.1e-70
Protein name				Locus	s Name	Acc#
				sp:TR	PG_PSEAE	P20576
Description						
TRANSFERASE)		<del>.</del>				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4145000_f2_47	1177	3097	415	1248	962	1.0e-96
Protein name				Locus	s Name	Acc#
Description				sp:00	P1_HAEIN	Q57242:005 056
ABC TRANSPORTER ATE	5 10 PX(115 P)	אר הססייביזאי	Υπγιο_1			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4181502_f3_149	1178	3098	130	393		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT			·			
	) THE T		NT	AA	0	Deck of 1 the
ORF Name	NTID	AAID	Length	Length	Score	Probability
4182762_f2_51	1179	3099	358	1077	374	1.9e-49
Protein name				Locu	s Name	Acc#
tryptophantRNA	ligase,			pir:H	70385	H70385
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
424203_t1_24	1180	3100	350	1053	542	3.2e-52
Protein name				Locus	s Name	Acc#
putative exodeox	yribonucle	ase (EC	3.1.11.2).	gp:SC	E87	AL132674
Description			- <del></del>			<del></del>
Streptomyces coe	licolor co	smid E87	•			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4331262_f3_141	1181	3101	222	669	402	2.2e-37
Protein name				Locus	s Name	Acc#
probable corA pr	otein			pir:F	70952	F70952
Description						
ORF Name	NTID	AAID	<u>NT</u> Length		Score	Probability
4506930_c1_174	1182	3102	356	1071	504	3.4e-48
Protein name				-	s Name XK HAEIN	Acc# P44491
Description					<del></del>	
TETRAACYLDISACCE	ARIDE 4'-F	(INASE, (	LIPID A 4'	-KINASE)		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4537837_£3_145	1183	3103	216	651	477	2.5e-45
Protein name					Name	Acc#
YciB homolog				gp:AF	114793	AF114793
Description				· · · · · · · · · · · · · · · · · · ·		
Vitreoscilla sp. outer membrane procomplete cds; and	otein, Bio	oA homol				_
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4571880_f1_16	1184	3104	216	651	437	4.3e-41
Protein name YbeZ protein					Name Y249116	<u>Acc#</u> AJ249116
<u>Description</u>						
Salmonella typhir	murium yle	eB (part:		ybez and	yber(pa	artial) genes.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4722125_c2_211	1185	3105	465	1398	1197	1.3e-121
Protein name  Description					Name 25_HAEIN	Acc# P44640
HYPOTHETICAL PRO	rein Hios:	25				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4798763_c1_155	1186	3106	141	426	274	8.1e-24
Protein name ExbD protein				_	Name E132741	Acc# AJ132741
Description						
Bordetella pertus ORF1(partial).	ssis hupB	, tonB, e	exbB, exbD	and bask	genes an	nd
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4876005_f1_30	1187	3107	828	2487	1231	3.1e-125
Protein name hypothetical prot	ein TM186	59		Locus	Name 72202	<u>Acc#</u> F72202
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4978375_c3_293	1188	3108	382	1149	261	7.7e-26
Protein name				Locus	s Name	Acc#
beta-ketoacyl-acyl	carrier	protein	synthase	pir:B	64545	B64545
Description				_		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5194068_f1_35	1189	3109	620	1863	1268	3.8e-129
<u>Protein name</u>					Name FX_HAEIN	Acc# P44933
Description						
ANTIPORTER)						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
5976555_£3_111	1190	3110	769	2310	3096	0.0
Protein name					Name	Acc#
Description						P00452:P78 088:P78177
(RIBONUCLEOTIDE RE	DUCTASE	1) (B1 P	ROTEIN) (R	1 PROTEIN	4)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
636513_f3_126	1191	3111	81	246		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6516_f2_45	1192	3112	361	1086	176	7.2e-13
Protein_name					Name	Acc#
hypothetical prote	ın			pir:S	76259	S76259
Description						

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Proba	bility
6828128_c2_223	1193	3113	201	606	105	0.000	66
Protein name phosphoglycerate mu	tase			Locus	s <u>Name</u> 72260		<u>Acc#</u> G72260
Description			·				
ORF Name	NTID	<u>AAID</u>	NT Length	AA Length [498	Score	Proba	bility
	1171	الــــــــا ا			L		
<u>Protein name</u>					s Name		<u>Acc#</u> Q55129
Description					-		~
HYPOTHETICAL 18.3 K	D PROTE	IN SLL040	0				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
786578_c3_259	1195	3115	86	261	99	2.8e-	05
Protein name				Locus	s Name		Acc#
unknown				gp:AF	114793		AF114793
Description				-			
Vitreoscilla sp. Youter membrane prote complete cds; and un	in, Bio	A homolog	tive tran , and glu	scription taminesyn	nal activ nthetase	ator,p homolo	outative og genes,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
969200_c3_270	1196	3116	234	705	360	6.2e-	33
Protein name					s Name DB_ECOLI		Acc# P17113
Description				L			
GLUCOSE INHIBITED D	IVISION	PROTEIN	В				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
970375_£2_67	1197	3117	279	840	921	2.2e-	92
Protein name			· · · · · ·	_	s <u>Name</u>		Acc#
probable GTP-bindin	g prote	ın H10393		pir:I	64150		I64150
Description							

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probal	oility
976531_£2_59	1198	3118	171	516	437	4.3e-4	1
Protein name YbeZ protein				Locus gp:STY2			Acc# AJ249116
Description							
Salmonella typhim	urium yl	eB (parti	al), miaB,	ybeZ and	ybeY (pa	irtial)	genes.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probal	oility
985925_f3_124	1199	3119	128	387			
<u>Protein name</u>				Locus	Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probab	oility
9954828_c2_208	1200	3120	322	969	170	3.9e-1	.4
Protein name TonB2				Locus gp:AF19			Acc# AF190125
Description				<u></u>			
Pseudomonas aerug complete cds.	inosa To	nB2 (tonB	32), ExbB (	exbB), and	ExbD (	(exbD) g	enes,
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probab	oility
10975667_c2_74	1201	3121	376	1131	524	2.6e-5	0
Protein name				Locus	Name		Acc#
thiamine-monophos	phate kir	nase		gp:D173	33		D17333
Description							
E. Coli thiL gene	, comple	e cds.					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probab	ility
12500081_c3_90	1202	3122	270	813	855	2.2e-8	5
Protein name				Locus sp:HIS6			Acc# P26721
Description					_		
HISF PROTEIN (CYC	LASE)						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
14572127_c1_50	1203	3123	165	498 433	1.1e-40
Protein name				Locus Name sp:RISB_ECOLI	Acc#
<u>Description</u>					P25540:P77
(LUMAZINE SYNTHA	SE) (RIBO	LAVIN SY	NTHASE BET	A CHAIN)	
ORF Name	NTID	AAID	NT Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
197212_c1_54	1204	3124	284	852	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
21767011_t3_32	1205	3125	317	954 736	8.9e-73
Protein name				Locus Name	Acc#
YafJ				gp:NGAJ2783	AJ002783
Description					
Neisseria gonorr	hoeae arol	(, aroB,	yaiJ genes	and open reading	frame.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23645263_c3_93	1206	3126	201	606 190	2.4e-30
Protein name				Locus Name	Acc#
				sp:PGPA_HAEIN	P44157
Description					
PHOSPHATIDYLGLYC	EROPHOS PHA	TASE A,			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23916007_c3_94	1207	3127	196	591 273	1.0e-23
Protein name	-			Locus Name	Acc#
methylase				gp:LLCPJW565	Y12736
Description					
Lactococcus lact	is cremor.	s plasmi	ld pJW565 D	NA, abiiM, abiiR	genesand orfX.

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23947167_£3_37	1208	3128	653	1962	756	6.0e-87
Protein name penicillin-binding	proteir	1 3		Locus pir:S	s Name 54872	Acc# S54872
Description						<del></del>
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24256687_£2_23	1209	3129	501	1506	537	1.8e-79
<u>Protein name</u>					s Name RF_ECOLI	Acc#
Description				<u> </u>		P11880:P77 636:O07100
(D-ALANYL-D-ALANIN	E-ADDING	ENZYME)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24353377_c1_45	1210	3130	225	678	155	3.3e-11
Protein name				Locus	s Name	Acc#
Protein name hypothetical protei	n PABOI	.31		Locus		Acc# D75209
	n PABOI	.31		_		<del></del>
hypothetical protein Description  ORF Name	NTID	AAID	NT Length	<u>AA</u> Length		<del></del>
hypothetical protein Description  ORF Name  26369016_f2_18				AA Length	75209 Score	D75209
hypothetical protein Description  ORF Name  26369016_f2_18  Protein name	NTID	AAID	Length	AA Length	75209	D75209
hypothetical protein Description  ORF Name  26369016_f2_18	NTID	AAID	Length	AA Length	75209 Score	D75209
hypothetical protein Description  ORF Name  26369016_f2_18  Protein name	NTID	AAID	Length	AA Length	75209 Score	D75209
hypothetical protein  Description  ORF Name  26369016_f2_18  Protein name  Description	NTID	AAID	Length	AA Length	75209 Score	D75209
hypothetical protein  Description  ORF Name  26369016_f2_18  Protein name  Description  NO-HIT	NTID 1211	<u>AAID</u>	Length 299 NT	AA Length 900 Locus	Score S Name	D75209  Probability  Acc#
hypothetical protein  Description  ORF Name  26369016_f2_18  Protein name  Description  NO-HIT  ORF Name	NTID 1211 NTID	AAID  AAID  AAID	Length 299  NT Length	AA Length  AA Length  AA Length  Locus  Locus  Locus	Score  Score Score	Probability  Acc#  Probability
hypothetical protein  Description  ORF Name  26369016_f2_18  Protein name  Description  NO-HIT  ORF Name  3397336_c1_49	NTID 1211 NTID	AAID  AAID  AAID	Length 299  NT Length	AA Length  AA Length  AA Length  Locus  Locus  Locus	Score Score Score Score Score	Probability  Acc#  Probability  0.037  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
35968792_f2_24	1213	3133	201	606	346	1.9e-31
Protein name					s Name	Acc# Q01893
Description						
TRIOSEPHOSPHATE	ISOMERASE,	(MIT)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3907500_£1_5	1214	3134	341	1026	227	9.2e-18
Protein name				_	s <u>Name</u>	Acc#
homoserine kinase	e nomolog			pir:T	33726	T33726
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3939665_c1_51	1215	3135	183	552	204	2.1e-16
Protein name					s Name SB HAEIN	Acc# P45150
Description						
N UTILIZATION SU	BSTANCE PR	ROTEIN B	HOMOLOG (N	USB PROTI	EIN)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3953191_c1_44	1216	3136	502	1509	1501	7.7e-154
Protein name glutamyI-tRNA syn	nthetase			_	Name 139107	Acc# AF139107
Description				] [32		
Pseudomonas aeru partial cds; hypo synthetase (gltX) complete sequence	thetical t genes, co	ranscrip	tional act	ivator(ad	ct) and g	lutamyl-tRNA
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
41703_f3_36	1217	3137	120	363		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT	<del></del>					

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4301943_f1_13	1218	3138	368	1107 1018	1.2e-102
Protein name				Locus Name sp:MRAY_HAEIN	Acc# P45062
Description					
(UDP-MURNAC-PENTA	PEPTIDE	PHOSPHOTRA	ANSFERASE)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5111318_£2_22	1219	3139	523	1572 756	6.8e-75
Protein name probable			<del>-</del>	Locus Name gp:AF141867	<u>Acc#</u> AF141867
Description		<u> </u>	*	<b>-</b>	
Vibrio cholerae probableUDP-N-acety ligase(murE) gene,			O-glutamat	e2,6-diaminopi	melate
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
6754052_f3_35	1220	3140	336	1011 695	2.0e-68
Protein name  Description				Locus Name sp:YABC_ECOLI	<u>Acc#</u> P18595
	VD - 5500		m HMCT TN	MEDAENTA DECTON	(ODED)
HYPOTHETICAL 34.9	KD PROT	EIN IN FRO			(ORFB)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
111552_f1_10	1221	3141	73	222	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
1189035_c3_48	1222	3142	179	540 616	4.7e-60
Protein name adenylate kinase				Locus Name gp:AB024426	Acc# AB024426
Description					
Pseudomonas putid	a adk ge	ne for ade	enylate ki	nase, complete co	ds.

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
12578208_f2_15	1223	3143	386	1161	1244	1.3e-	126
Protein name					s Name AS_PSEAE		<u>Acc#</u> Q51344
Description					-		
DEHYDROGENASE)							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
23444507_c3_45	1224	3144	452	1359			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
23536511_f2_16	1225	3145	338	1017	210	4.1e-	15
Protein name					s Name G1 ECOLI		<u>Acc#</u> P18840
Description				L			`
(L-ASNASE I)						_	
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Proba	bility
245682_f1_8	1226	3146	303	912	689	8.5e-	68
Protein name					s Name UA ECOLI		Acc# P07649
Description							
1) (PSEUDOURIDINE	SYNTHASI	E I) (URAC	IL HYDROL	YASE) (P	SU-I)	<u></u>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
34157662_c2_41	1227	3147	203	612	321	8.5e-	29
<u>Protein name</u>					s Name PB_PSEFL		<u>Acc#</u> P52237
Description							
BIOGENESIS PROTEIN	TIPB)					<del>-</del> -	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4042131_c1_33	1228	3148	70	213	
Protein name				Locus Name	Acc#
Description					
NO-HIT		<del></del>			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
4112793_c1_37	1229	3149	423	1272 175	3.1e-10
Protein name				Locus Name sp:CCMH_HAEIN	Acc# P46458
Description				-	<del></del>
CYTOCHROME C-TYPE	BIOGENE	SIS PROTE	IN CCMH PR	ECURSOR	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4484436_c2_40	1230	3150	692	2079 1743	1.7e-179
Protein name				Locus Name sp:CCMF_PSEFL	Acc# P52225
Description					<del>,,,,,</del>
CYTOCHROME C-TYPE	BIOGENE	SIS PROTE	IN CYCK		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5265800_£1_9	1231	3151	77	234 274	8.1e-24
Protein name				Locus Name	Acc# P20458
Description					
TRANSLATION INITI	ATION FA	CTOR IF-1			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
587775_c1_36	1232	3152	172	519 299	1.8e-26
Protein name				Locus Name sp:CCMH_ECOLI	Acc# P33925
Description				<u>.</u>	

ORF Name	NTID AAID	<u>NT</u> Length	AA Score	Probability
5894082_t1_7	1233 3153	206	621 309	1.6e-27
Protein name			Locus Name sp:YHHF_ECOLI	Acc# P10120
Description				<del>,</del>
21.7 KD PROTEIN IN	FTSY-NIKA INTERG	ENIC REGI	ON	· · · ·
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
1058462_c3_105	1234 3154	283	852 77	0.032
Protein name			Locus Name	Acc#
15 kDa vesicular-l	ike antigen		gp:PFAVLAP	M94732
Description				
Plasmodium falcipa	rum 15 kDa vesicu	lar-like	antigen gene, exc	ons 1through 4.
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
13688802_c2_101	1235 3155	76	231	
Protein name			Locus Name	Acc#
Description				
NO-HIT				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
14644586_f2_28	1236 3156	391	1176 465	4.7e-44
Protein name  36 kDa protein			Locus Name gp:HPU86610	<u>Acc#</u> U86610
Description			J [55	
Helicobacter pylor	i 36 kDa protein (	gene, com	plete cds.	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
16132787_±3_50	1237 3157	105	318 197	1.2e-15
Protein name			Locus Name sp:YDCQ_ECOLI	<u>Acc#</u> P76107
Description				
HYPOTHETICAL 16.1	KD PROTEIN IN TEH	B-ANSP IN	TERGENIC REGION	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19532661_£2_33	1238	3158	77	234		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT	· · · · · · · · · · · · · · · · · · ·					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20335427_c2_78	1239	3159	657	1974	239	2.0e-17
Protein name minor tail protein	n gp26-re	elated prot	ein	Locus	5 Name 75605	Acc# F75605
Description	<del></del>			J [		
ORF Name	NTID	<u>AAID</u>	NT Length	AA Length	Score	Probability
Protein name			L	Locus	s Name	Acc#
Description				2004.	<u> </u>	120.517
NO-HIT						
			NT	AA		
ORF Name	NTID	AAID	Length	<u>Length</u>	Score	Probability
21909377_f2_29	1241	3161	401	1206	261	3.0e-21
Protein name				Locus	s Name	Acc#
hypothetical prote	ein jhp13	380		pir:G	71815	G71815
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22266577_£2_21	1242	3162	220	663	233	1.8e-19
Protein name				Locus	Name	Acc#
thiamine-phosphate	e pyropho	sphorylase	;	gp:AF	180145	AF180145
Description				- L.		<del></del>
Zymomonas mobilis protein yidC (yidC gltB (gltB), glutan udk (udk), hypothet protein; zml2orf5, A, beta-hydroxyster	), hypoth mate synt ical prot hypothet:	netical pro chasesmall cein, NADH ical prote:	otein,glu subunit dehydrog in, aspar	tamine-pygltS (glt enase, hy tate amin	yruvate a (S), unde pothetic notransfe	aminotransferase ecaprenol kinase cal

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
22894061_c3_119	1243	3163	165	498 222	2.6e-18
Protein name				Locus Name sp:TOLR_PSEAE	Acc# P50599
Description					
TOLR PROTEIN					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23444426_f3_45	1244	3164	419	1260 642	8.3e-94
Protein name				Locus Name	Acc#
ATP-dependent hel	icase Hr	A homolo	g.	gp:D90779	D00770.D00
<u>Description</u>					D90779:D90 761:AB0013 40
E.coli genomic DN	A, Kohara	a clone #	268 (31.6-3	2.0 min.).	· · · · · ·
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24021016_f3_43	1245	3165	1195	3588 2225	1.6e-266
Protein name	•			Locus Name	Acc# P45128
Description					······································
TRANSCRIPTION-REP	AIR COUP	LING FACT	OR (TRCF)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24401887_c1_62	1246	3166	114	345	2.9e-12
Protein name				Locus Name gp:AB030825	Acc# AB030825
Description					
Pseudomonas aerug	inosa ge	nomic DNA	, partial	sequence, strain	: PAO1 .
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
25554561_t2_20	1247	3167	151	456 94	0.0015
Protein name	oin Duio	\1		Locus Name	Acc#
hypothetical prot	ern burn	) <u> </u>		pir:D71092	D71092
Description					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2994032_c2_82	1248	3168	264	795	307	9.6e-35
Protein name				Locu	s Name	Acc#
minor tail protein	gp19			pir:T	13105	T13105
Description				<u>.</u>		
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
31916632_f1_8	1249	3169	62	189		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT					<del> </del>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3320327_c2_76	1250	3170	94	285		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34188892_c1_61	1251	3171	673	2022	227	4.5e-15
Protein name					s Name 26_BPMD2	Acc# 064220
Description						
MINOR TAIL PROTEIN	GP26			<u> </u>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34415711_£1_10	1252	3172	368	1107	288	2.7e-25
Protein name				Locus	s Name	Acc#
conserved hypothet protein HP1486	ical in	tegral meml	orane	pir:F	64705	F64705
Description				_		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35367058_f3_51	1253	3173	76	231	151	8.8e-11
Protein name					s Name CQ_ECOLI	<u>Acc#</u> P76107
Description						
HYPOTHETICAL 16.1	KD PROT	EIN IN TEH	B-ANSP IN	TERGENIC	REGION	
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
35942905_£2_19	1254	3174	156	471	278	3.1e-24
Protein name					s Name BK_ECOLI	Acc# P33899
Description						
HYPOTHETICAL TRNA	RRNA ME	THYLTRANSF	ERASE YIE	BK,		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36118750_c2_104	1255	3175	77	234		
Protein name				Locu	s_Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36328956_t2_23	1256	3176	108	327		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3944450_c2_93	1257	3177	232	699	422	1.7e-39
Protein name TolQ protein	· · · · · ·			Locus	S Name	Acc# X74218
Description	·····					
Pseudomonas putida	ruvB.	tolO, tolR	, tolA, t	olB and	oprL gene	s.

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4460387_t1_9	1258	3178	514	1545	240	3.4e-17
Protein name hypothetical prote	in jhpl	882		Locus pir:A	3 Name 71816	Acc# A71816
Description				<u> </u>		
ORF Name 4507703 c2 103	NTID 1259	<u>AAID</u>	NT Length	AA Length 309	Score	Probability 2.0e-11
Protein name	J [		L		Name 14_BPHP1	Acc# P51716
Description				<u> </u>		
HYPOTHETICAL 14.9	KD PROTI	EIN IN REP	-HOL INTE	RGENIC RI	EGION (OF	RF14)
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
4728415_c3_120	1260	3180	276	831	103	0.030
Protein name ras interacting pro	otein Rl	PA		_	Name 159241	Acc# AF159241
<u>Description</u>				J <u>[</u>		
Dictyostelium disco	oideum i	ras intera	cting pro	tein RIPA	A (ripA)	mRNA, complete
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
4730050_c1_73	1261	3181	439	1320	375	1.6e-34
Protein name TolB					Name U32470	Acc# U32470
Description						<del></del>
Haemophilus influer gene, partial cds, protein (tolA) and (	inner me	embrane pro	otein (to	lR), out	ermembran	ne integrity
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5282805_c1_63	1262	3182	227	684	329	1.2e-29
Protein name				Locus	Name	Acc#
minor tail protein	L homol	.og:protei	gp18	pir:T	13104	T13104
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5348393_c2_83	1263	3183	79	240	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
682777_c2_79	1264	3184	139	420	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
7265950_f1_7	1265	3185	1014	3045 726	3.1e-134
Protein name				Locus Name sp:HRPA_ECOLI	
<u>Description</u>				•	P43329:P77 479:P76861 :P76863
ATP-DEPENDENT HE	LICASE HR	PA			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24113927_f2_1					
	1266	3186	334	1005	0.0012
Protein name	1266	3186	334	Locus Name	0.0012 Acc#
STARP antigen	1266	3186	334	1005 116	
	[1266	3186	334	Locus Name	Acc#
STARP antigen				Locus Name	Acc#
STARP antigen  Description				Locus Name	Acc#
STARP antigen  Description  P.reichenowi STA	RP gene fo	or STARP	antigen.	Locus Name  gp:PRSTARPA  AA  Score	Acc# Z30339
STARP antigen  Description  P.reichenowi STA  ORF Name	RP gene fo	or STARP	antigen.  NT Length	Locus Name  gp:PRSTARPA  AA Length 621 Locus Name	Acc# Z30339  Probability 7.0e-41  Acc#
STARP antigen  Description  P.reichenowi STA  ORF Name  25673906_f1_1	RP gene fo	or STARP	antigen.  NT Length	Locus Name  gp:PRSTARPA  AA Length  621  435	Acc# Z30339  Probability 7.0e-41  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29927207_f3_4	1268	3188	230	693	88	2.4e-05
Protein name probable two compon	ient sei	nsor pro	tein	Locus pir:C	Name 70624	Acc# C70624
Description				<u> </u>		
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
35210305_f1_2	1269	3189	300	903	155	2.3e-08
Protein name					Name	Acc#
SmeS				gp:AF	173226	AF173226
Description						
Stenotrophomonas ma (smeS), SmeA (smeA),	Iltophi SmeB	lia mult (smeB),	idrug efflu and SmeC (s	x system smeC) gene	SmeR (sr es,comple	neR),SmeS ete cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12938586_c3_89	1270	3190	152	459	296	3.8e-26
Protein name		_ ·			Name	Acc# P43036
Protein name  Description						
	CIATED	LIPOPRO1	EIN PRECURS	sp:PA		
Description	CIATED :	LIPOPROT AAID	EIN PRECURS  NT  Length	sp:PA		
Description PEPTIDOGLYCAN-ASSOC			NT	SOR AA	L_PSEPU	P43036
Description PEPTIDOGLYCAN-ASSOCO	NTID	AAID	<u>NT</u> Length	SP:PA  SOR  AA  Length  258	L_PSEPU	P43036
Description  PEPTIDOGLYCAN-ASSOC  ORF Name  [14237656_c1_39]	NTID	AAID	<u>NT</u> Length	SP:PA  SOR  AA  Length  258	L_PSEPU Score	P43036 Probability
Description  PEPTIDOGLYCAN-ASSOC  ORF Name  14237656_c1_39  Protein name	NTID	AAID	<u>NT</u> Length	SP:PA  SOR  AA  Length  258	L_PSEPU Score	P43036 Probability
Description  PEPTIDOGLYCAN-ASSOC  ORF Name  14237656_c1_39  Protein name  Description	NTID	AAID	<u>NT</u> Length	SP:PA  SOR  AA  Length  258	L_PSEPU Score	P43036 Probability
Description  PEPTIDOGLYCAN-ASSOC  ORF Name  14237656_c1_39  Protein name  Description  NO-HIT	NTID 1271	<u>AAID</u>	NT Length 85 NT	SOR  AA  Length  258  Locus	Score S Name	P43036  Probability  Acc#
Description  PEPTIDOGLYCAN-ASSOC  ORF Name  14237656_c1_39  Protein name  Description  NO-HIT  ORF Name	NTID 1271 NTID	AAID  AAID	NT Length 85 NT Length	Sp:PA  SOR  AA  Length  Locus  AA  Length  831	Score S Name	P43036  Probability  Acc#
Description  PEPTIDOGLYCAN-ASSOC  ORF Name  14237656_c1_39  Protein name  Description  NO-HIT  ORF Name  14492157_f3_31	NTID 1271 NTID	AAID  AAID	NT Length 85 NT Length	Sp:PA  SOR  AA  Length  Locus  AA  Length  831	Score Score Score	Probability  Acc#  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14875327_f2_18	1273	3193	867	2604	1592	1.7e-163
Protein name  membrane alanyl a	minopepti	Ldase			s Name 157493	Acc# AF157493
Description						
Zymomonas mobilis	ZM4 fost	mid clone	42D7, com	plete se	quence.	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
156258_c3_90	1274	3194	199	600	126	2.6e-06
Protein name NrpG					s <u>Name</u> U46488	Acc# U46488
Description						
Proteus mirabilis NrpA (nrpA), NrpB	_		-	_	_	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16180387_£3_36	1275	3195	384	1155	143	3.8e-07
Protein name				Locus	s Name	Acc#
hypothetical prot	ein RP367	7		<b>□</b> □ □ □ □ ▼ ▼	71602	H71693
				pir:H	71093	H/1693
Description		·		] рт:н	71693	M/1693
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
ORF Name	NTID	AAID	Length	AA Length 1017 Locus	Score	Probability
ORF Name 16507676_c2_64	NTID	AAID	Length	AA Length 1017 Locus	Score 445 S Name	Probability 6.1e-42
ORF Name  16507676_c2_64  Protein name	NTID	AAID	Length	AA Length 1017 Locus	Score 445 S Name	Probability  6.1e-42  Acc#  P36566:P77
ORF Name  16507676_c2_64  Protein name  Description	NTID	AAID	Length	AA Length 1017 Locus	Score 445 S Name	Probability  6.1e-42  Acc#  P36566:P77
ORF Name  16507676_c2_64  Protein name  Description  SMTA PROTEIN	NTID 1276	<u>AAID</u> 3196	Length  338  NT	AA Length 1017 Locus Sp:SM	Score 445 S Name TA_ECOLI	Probability 6.1e-42 Acc# P36566:P77 586
ORF Name  16507676_c2_64  Protein name  Description  SMTA PROTEIN  ORF Name	NTID 1276 NTID	AAID  AAID	Length  338  NT  Length	AA Length  Locus Sp:SM  AA Length  330 Locus	Score  445  Name TA_ECOLI  Score  205  Name	Probability  6.1e-42  Acc#  P36566:P77 586  Probability  1.7e-16  Acc#
ORF Name  16507676_c2_64  Protein name  Description  SMTA PROTEIN  ORF Name  22116326_t2_14	NTID 1276 NTID	AAID  AAID	Length  338  NT  Length	AA Length  Locus Sp:SM  AA Length  330 Locus	Score  445  Name  TA_ECOLI  Score	Probability  6.1e-42  Acc#  P36566:P77 586  Probability  1.7e-16

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
22694056_f1_11	1278	3198	637	1914 1865	2.1e-192
Protein name				Locus Name sp:CLPB_HAEIN	Acc# P44403
Description					
CLPB PROTEIN					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23495633_t2_15	1279	3199	349	1050 105	0.0072
Protein name				Locus Name	Acc#
ComB				gp:AF027189	AF027189
Description					
Acinetobacter sp. and unknown genes.	BD413 I	ytB, comB,	comC, co	mE, and comF gene	es,complete cds;
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Length Score	Probability
2376890_c2_56	1280	3200	90	273	
Protein name				Locus Name	Acc#
Description	•				
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24395640_f3_38	1281	3201	282	849 291	1.3e-25
Protein name				Locus Name	Acc#
ABC transporter po	tG			pir:B71694	B71694
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24643831_f1_3	1282	3202	346	1041 213	3.2e-16
Protein name				Locus Name	Acc#
phospholipase A				gp:CCPLDA	Y11031
Description				_	
C.coli pldA gene.					

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24783453_f2_25	1283 3203	224	675	738	5.5e-73
Protein name				s <u>Name</u> PB_BACNO	Acc# P17422
Description					
CLPB PROTEIN					
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25817157_£3_34	1284 3204	250	753	319	1.4e-28
Protein name hypothetical protei	n			Name WAAA179	Acc# Z96927
Description					
Acinetobacter haemo	olyticus waaA gene	e, strain	ATCC 17	906.	<u></u>
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2995252_f3_37	1285 3205	342	1029	198	1.4e-13
Protein name ct391 hypothetical	nrotoin		_	s Name	Acc#
	procein		pir:G	72072	G72072
Description					
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
32703126_f2_22	1286 3206	304	915	369	6.9e-34
Protein name			Locus	Name	Acc#
hypothetical protei	n RP368		pir:A	71694	A71694
Description					
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35368941_f1_4	1287 3207	285	858	152	4.5e-09
Protein name competence protein	ComF			Name 1249742	Acc# AJ249742
Description			J <u>Dr. 1</u>		
Pseudomonas stutzer	ri JM300 bioB (par	ctial), c	omF and o	of (part	ial)genes.

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35943885_c2_66	1288	3208	413	1242		
Protein name				Locu	s Name	Acc#
Description			,			
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4111633_f2_13	1289	3209	154	465		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4142186_f1_2	1290	3210	246	741	777	4.0e-77
Protein name					s Name	<u>Acc#</u>
Description				sp:RN	PH_PSEAE	P50597
NUCLEOTIDYLTRANSFI	ERASE)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4572203_f1_9	1291	3211	329	990	117	5.7e-05
Protein name		_			s Name	Acc#
merozoite surface	antigen	2		gp:09	1655	U91655
Description						
Plasmodium falcipa partial cds.	arum iso	late V310	, merozoit	e surface	e antiger	n 2(MSP-2) gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4797282_c2_74	1 13 7 7 7 T	3212	67	204		
•	1292			204		
Protein name	1292	_] [3212	] []	ļ	s Name	Acc#
Protein name Description			J []	ļ	s Name	Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5082637_£3_33	1293	3213	440	1323	635	4.5e-62
Protein name WaaA					s Name 026386	Acc# AF026386
Description						
Salmonella typhim region, WaaY (waaY WaaP (waaP), WaaG gene, partial cds.	) gene, ]	partial (	cds; WaaJ (	(waaJ),Wa	aI (waaI)	), WaaB (waaB),
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5132838_f1_5	1294	3214	255	768		
Protein name  Description				Locu	s Name	Acc#
NO-HIT						
ORF Name 5867075_f3_29	NTID 1295	AAID 3215	NT Length	AA Length	Score 105	Probability 0.00049
Protein name pilV protein				Locu pir:S	s <u>Name</u> 77594	<u>Acc#</u> S77594
Description						
ORF Name 790700_c2_65	<u>NTID</u>	<u>AAID</u>	NT Length	AA Length 1140	Score 151	Probability 9.2e-08
Protein name				Locu	s Name	Acc#
hypothetical prote	ein TP056	55		pir:C	71308	C71308
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
9775283_c1_46	1297	3217	499	1500	469	1.8e-44
Protein name probable alginate (algI) Description	O-acety]	ation pr	rotein	Locus pir:D	71308	Acc# D71308

ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
115827_c1_7	1298	3218	330	993	877	1.0e-87
Protein name					s Name MU_HAEIN	Acc# P43889
Description					-	<del></del>
ACETYLGLUCOSAMINE-1	-PHOSPH	ATE URIDY	CTRANSFER	ASE)		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16828178_f1_2	1299	3219	616	1851	2166	2.6e-224
Protein name				-	S Name PA_HAEIN	Acc# P44910
Description						
GTP-BINDING PROTEIN	TYPA/E	IPA HOMOLO	OG			
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
32656465_c2_10	1300	3220	78	237		
<u>Protein name</u>				Locus	s Name	Acc#
Description						
NO-HIT						***
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3336053_f1_1	1301	3221	133	402	618	2.9e-60
Protein name				Locus	Name	Acc#
outer membrane prot	ein CD	precursor		pir:S	39866	S39866
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10975831_c3_12	1302	3222	92	279		
Protein name				Locus	Name	Acc#
Description						
NO-HIT		<u> </u>				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
15912757_c1_8	1303	3223	123	372	86	0.048
Protein name FIP2				Locus N gp:AF061		Acc# AF061034
Description						
Homo sapiens FI	2 alterna	tively t	ranslated m	nRNA, comple	te cds	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
22457187_f3_5	1304	3224	250	753	388	7.0e-89
Protein name				Locus N sp:Y882		Acc# P44068
Description						
HYPOTHETICAL PRO	TEIN HI08	82				
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>Length</u>	ore	Probability
35271883_f1_4	1305	3225	60	183		
Protein name				Locus N	ame	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
22848457_f2_3	1306	3226	134	405		
Protein name			•	Locus N	ame	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
22853376_f3_5	1307	3227	239	720		
Protein name				Locus N	ame	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
29296968_c3_9	1308	3228	77	234	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
976678_c3_10	1309	3229	190	570 265	7.3e-23
Protein name				Locus Name sp:PRTR_PSEA	Acc# Q06553
Description					
TRANSCRIPTION REC	JULATORY	PROTEIN PE	RTR (PYOSI	N REPRESSOR PRO	TEIN)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
989077_£1_1	1310	3230	122	369	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
1062575_c3_34	1311	3231	109	330	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
11125280_£3_13	1312	3232	146	538	8.6e-52
Protein name	olog HI03	77		Locus Name	Acc# C64064
Description					

ORF Name	NTID AAID Length	Score	Probability
1297092_c1_20	1313 3233 111	336 174	3.9e-12
Protein name  probable gamma-glut precursor	camyItranspeptidase	Locus Name	<u>Acc#</u> E70682
Description	- 10	<del></del>	
ORF Name	NTID AAID No. 100 Leng	- Score	Probability
Protein name		Locus Name	Acc#
probable gamma-glut	tamyltranspeptidase	pir:T34901	T34901
Description			
ORF Name 20727194 c2 24	NTID AAID No. 1	– Score	Probability
Protein name		Locus Name  [gp:AF017750	Acc# AF017750
Description			
(ccmH), recombination	cytochrome C-type binal DNA repair protein (sodA), and CitG prot	(recR), manganese	enes,
ORF Name	NTID AAID NTID Leng	- — Score	Probability
21679025_±1_1	] [1316   3236   420	1263 1516	2.0e-155
Protein name	· · · · · · · · · · · · · · · · · · ·	I agus Nome	
		Locus Name sp:NIFS ECOLI	<u>Acc#</u>
Description		sp:NIFS_ECOLI	Acc# P39171:P76 581:P76992
	DG .		P39171:P76
Description	OG NTID AAID Leng	sp:NIFS_ECOLI  AA Score	P39171:P76
Description  NIFS PROTEIN HOMOLO	NTID AAID NI	sp:NIFS_ECOLI	P39171:P76 581:P76992
Description  NIFS PROTEIN HOMOLO  ORF Name	NTID AAID NTID Leng	sp:NIFS_ECOLI  F AA Score  Gth Length	P39171:P76 581:P76992 Probability
Description  NIFS PROTEIN HOMOLO  ORF Name  31797152_f3_14	NTID AAID NTID Leng	sp:NIFS_ECOLI  F AA Score  gth Length 224  Locus Name	P39171:P76 581:P76992 Probability 1.6e-18 Acc#

ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
32244203_c2_26	1318	3238	72	219	125	5.0e-08
Protein name				-	Name H231122	Acc# AJ231122
Description						
Vibrio cholerae z61	.f gene					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33398287_t2_5	1319	3239	178	537	396	9.6e-37
Protein name					Name HP_HAEIN	Acc# P44675
Description				<u></u>		<del></del>
HYPOTHETICAL PROTEI	N HIO3	79				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36129678_f1_2	1320	3240	112	339	384	1.8e-35
Protein name					Name HF_HAEIN	Acc# P44672
Description						
HYPOTHETICAL PROTEI	N HIO3	76				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
36220382_c2_25	1321	3241	120	363	174	3.2e-12
<u>Protein name</u>				Locus	Name	Acc# P20735
Description						
GLUTAMYLTRANSFERASE	(GGT)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4331938_f2_9	1322	3242	622	1869	1435	7.6e-147
Protein name					Name	Acc# P44669
Description				i		
CHAPERONE PROTEIN H	SCA (HS	C66)			·	

ORF Name	NTID AAID NT Length	AA Score	Probability
4332838_£2_10	1323 3243 115	348 397	7.5e-37
Protein name ferredoxin		Locus Name	Acc# AF096864
Description			
_	inosa heat shock protein ( (fdx), and nucleoside dip		
ORF Name 5898593_c2_28	NTID AAID NT Length	AA Score Length	Probability
Protein name Description		Locus Name	<u>Acc#</u>
NO-HIT			
ORF Name	NTID AAID NT Length	<u>AA</u> <u>Length</u> <u>Score</u>   486   351	Probability 5.6e-32
Protein name  putative gamma-gluprecursor	utamyItranspeptidase	Locus Name gp:PST249741	Acc# AJ249741
Description		<u> </u>	
Pseudomonas stutz	eri JM300 gacS (partial) a	nd ggtB (partial)	genes.
ORF Name 12915808_f3_10	NTID AAID NT Length	AA Score	Probability
Protein name		Locus Name	Acc#
Description			
NO-HIT			
ORF Name	NTID AAID <u>NT</u> Length	AA Length Score	Probability
20737503_£3_8	3247 371	1116 418	4.5e-39
Protein name probable permease	perM homolog (perM) RP630	Locus Name	<u>Acc#</u> E71668
Description			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22000293_c2_13	1328	3248	97	294	348	1.2e-31
Protein name 50S ribosomal prot	ein homo	olog			s Name 153712	Acc# AF153712
Description						
Pseudomonas sp. Bo complete cds.	33R str	ain BG33R	50S ribos	omal pro	tein homo	ologgene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23863307_f3_9	1329	3249	261	786	194	2.4e-15
Protein name		·			s Name GE_HAEIN	<u>Acc#</u> 086235
Description						
HYPOTHETICAL PROTI	EIN HI12	25.1				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24308561_c3_17	1330	3250	182	549	710	5.1e-70
Protein name				Locu	s Name	Acc#
phosphoribosylform cyclo-ligase,:5'-ph ole synthetase			ninoimidaz	pir:A	JECPC	A25955:B65
Description				_		
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
26277251_c3_18	1331	3251	131	396	352	4.4e-32
Protein name					s <u>Name</u> R5_ECOLI	Acc# P08178
Description						
(PHOSPHORIBOSYL-AN	MINOIMID	AZOLE SYNT	THETASE) (	AIR SYNT	HASE)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6142915_c2_14	1332	3252	228	687	382	2.9e-35
Protein name				Locus	s Name	Acc#
5'-phosphoribosylo	lycinam:	ide transf	ormylase	gp:ST	068765	U68765
Description				-		
Salmonella typhimu 5'-phosphoribosyl-						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10744000_c3_102	1333	3253	309	930	1094	1.0e-110
Protein name probable Mn trans	port prot	ein		Locus	8 Name 64063	Acc#
Description				J [		G64063:C41 833
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
1181631_£1_2	1334	3254	558	1677	1333	4.9e-136
<u>Protein name</u>					s Name IM_PSEPU	<u>Acc#</u> P25754
Description						
60 KD INNER-MEMBR	ANE PROTE	EIN				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13703378_c3_117	1335	3255	95	288	163	4.7e-12
Protein name					Name AQ_ECOLI	Acc# P76246
Description						····
HYPOTHETICAL 8.7	KD PROTE	N IN GAPA	-RND INTE	RGENIC RI	EGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
15031513_f3_43	1336	3256	479	1440	1390	4.5e-142
Protein name					Name RC_METGL	Acc# P37145
Description				<u> </u>		
THREONINE SYNTHAS	Ξ,					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15039077_c1_64	1337	3257	266	801	196	1.5e-15
Protein name					Name INTREG	Acc# X98546
Description				OF 1		
D.nodosus intB, re	egA, gepA	, gepB, ar	nd gepC g	enes.		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
15665903_c1_59	1338	3258	281	846 1074	1.4e-108
Protein name				Locus Name sp:Y360_HAEIN	Acc# P44661
Description					
HYPOTHETICAL PROTE	IN HIO3	50			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
15798825_c1_65	1339	3259	165	498	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19538327_c1_72	1340	3260	213	642 218	7.0e-18
Protein name	-			Locus Name	Acc#
				sp:Y882_METJA	Q58292
Description					
HYPOTHETICAL PROTE	SOUM NI	32			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
197188_f3_42	1341	3261	345	1038 659	1.3e-64
Protein name				Locus Name	Acc#
				sp:FMT_PSEAE	085732
Description					
METHIONYL-TRNA FOR	MYLTRAN	SFERASE,			11.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20197175_f3_44	1342	3262	415	1248 453	5.0e-47
Protein name				Locus Name	Acc#
				sp:SMF_HAEIN	P43862
Description					
SMF PROTEIN (DNA P	ROCESSI	NG CHAIN A	<u> </u>	<del>-</del>	

ORF Name	NTID A	AID	<u>NT</u> Length	AA Length	Score	Probability
23440886_f2_27	1343	3263	602	1809	241	2.1e-19
<u>Protein name</u>				-	Name 78_METJA	<u>Acc#</u> Q58091
Description				-	<del>-</del> -	
HYPOTHETICAL PROTEI	N MJ0678			<del>.</del>		
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
241290_f2_31	1344	3264	65	198		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						-
ORF Name	NTID AF	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24244010_c3_106	1345	265	83	252	69	0.042
Protein name				Locus	Name	Acc#
hypothetical protei	n Y105C5B.	х		pir:T2	26400	T26400
Description						
ORF Name	NTID AF	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24253187_c3_107	1346	266	66	201		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID AF	<u> </u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24255262_c2_96	1347	267	430	1293	873	2.7e-87
Protein name				Locus	Name	Acc#
conserved hypotheti	cal protei	n		pir:C7	75339	C75339
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24256550_f3_40	1348	3268	165	498	465	4.7e-44
Protein name					s Name AD_ECOLI	Acc# P25538
Description						
HYPOTHETICAL 17.2	KD PROTI	EIN IN TSX	-RIBG INT	ERGENIC	REGION (C	ORF1)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24337786_f3_48	1349	3269	311	936	657	2.1e-64
Protein name				•	s Name GI_BRUAB	<u>Acc#</u> Q59174
Description						
ARGINASE,						
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24417762_f1_18	1350	3270	62	189	74	0.030
Protein name					s Name 1A_SERMA	<u>Acc#</u> P22595
Description				L		
TYPE-1 FIMBRIAL PRO	OTEIN ST	DBUNIT PRE	CURSOR			
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24489626_f2_21	1351	3271	474	1425	1058	6.8e-107
Protein name					s Name DF_PSEPU	Acc# P25755
Description						
POSSIBLE THIOPHENE	AND FU	RAN OXIDAT	ION PROTE	IN THDF		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24643777_f2_22	1352	3272	352	1059	682	4.3e-78
Protein name					s Name BD ECOLI	Acc# P25539
Description					_	
RIBOFLAVIN-SPECIFIC	C DEAMII	NASE,	-			

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
24643937_f1_4	1353	3273	450	1353 695	2.0e-68
Protein name				Locus Name	Acc# P44788
Description					<del></del>
SUN PROTEIN (FMU P	ROTEIN)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
26603562_c2_86	1354	3274	303	912 1047	9.9e-106
Protein name				Locus Name sp:FECE_HAEIN	Acc# P44662
Description					
IRON(III) DICITRATE	E TRANS	PORT ATP-	BINDING PR	OTEIN FECE HOMOLO	OG
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
2738783_f2_37	1355	3275	60	183	
Protein name				Locus Name	Acc#
Description					
NO-HIT			_		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
2750262_±1_1	1356	3276	103	312 193	3.1e-15
Protein name hypothetical protein	n SCH24	1.04		Locus Name	Acc# T36569
Description					
ORF Name	NTID	AAID	NT Length	AA Score	Probability
29539015_c1_62	1357	3277	417	1254 666	2.3e-65
Protein name				Locus Name sp:YDHH_ECOLI	Acc# P77570
Description					<u> </u>
HYPOTHETICAL 39.5	KD PROT	EIN IN PD	XH-SLYB IN	TERGENIC REGION	

ORF Name	NTID	AAID	<u>NT</u> Length	Length Score	Probability
30739700_£1_7	1358	3278	211	636 244	1.2e-20
Protein name				Locus Name sp:YRDC_ECOLI	Acc# P45748
Description					
HYPOTHETICAL 20.	8 KD PROT	EIN IN ARC	E-SMG INT	ERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
34088143_f3_55	1359	3279	106	321	
Protein name				Locus Name	Acc#
Description					
NO-HIT			<del>-</del>		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
3913215_f2_26	1360	3280	165	498	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3939063_f2_23	1361	3281	225	678 519	8.8e-50
Protein name				Locus Name sp:RISA_PHOPO	Acc# P51961
Description					
RIBOFLAVIN SYNTH	ASE ALPHA	CHAIN,			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3942213_c2_97	1362	3282	367	1104 930	2.5e-93
Protein name				Locus Name sp:GCH2_PHOLE	Acc# Q02008
Description					

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
4785911_t2_29	1363	3283	435	1308	9.1e-135
Protein name				Locus Name sp:OAT_DROAN	Acc# P49724
Description					
ACID AMINOTRANSFERA	ASE)				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5214052_f3_53	1364	3284	411	1236 1096	6.4e-111
Protein name				Locus Name	Acc# P43836
Description					
TYROSYL-TRNA SYNTHE	ETASE,	(TYROSINE-	-TRNA LIG	ASE) (TYRRS)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
5282562_c1_63	1365	3285	265	798 693	3.2e-68
Protein name				Locus Name	Acc#
hypothetical protei	n jhp0:	330		pir:B71947	B71947
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
6070166_f2_20	1366	3286	71	216	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
6147028_c1_60	1367	3287	292	879 870	5.6e-87
Protein name				Locus Name sp:YFED_YERPE	<u>Acc#</u> Q56955
Description				1	
CHELATED IRON TRANS	PORT S	YSTEM MEMB	RANE PROT	EIN YFED	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
839752_f1_19	1368	3288	60	183			
Protein name				Locus	s Name		Acc#
Description							
по-ніт							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
867183_c1_68	1369	3289	128	387	107	2.7e-0	05
Protein name					Name  AM_BACSU		<u>Acc#</u> 007931
Description				<u> </u>			
HYPOTHETICAL 39.5	KD PROTEI	N IN SIGZ	-CSN INT	ERGENIC P	REGION	<del></del>	
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probab	oility
1197077_f3_44	1370	3290	375	1128	178	8.2e-1	.1
Protein name	_			Locus	s Name		Acc#
hypothetical prote	in TM0342			pir:D	72388		D72388
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
14641008_f3_46	1371	3291	271	816	360	6.2e-3	3
Protein name  putative thiol:dis	ulfide in	terchange	protein		Name 057031		Acc# AF057031
Description				] [9p.m.			AL 03 / 03 L
Pseudomonas aerugi	noga nufa	tive thic	ol dienif	ide inter	rchange r	rotein	recursor
(dsbC) gene, comple	_						JI COULDOI
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
15058126_f1_9	1372	3292	204	615	183	3.6e-1	4
Protein name hypothetical prote	<u>in</u>			_	Name 088857		<u>Acc#</u> AF088857
Description				٠٠٠٠٠ [ع			-11 00000,
Vogesella indigofe sequence.	ra indigo	idine bio	synthesi	s regulat	cory locu	s,comp	Lete

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
158638_c3_82	1373	3293	89	270 350	7.2e-32
Protein name				Locus Name sp:IMDH_ACICA	Acc# P31002
Description					·
DEHYDROGENASE) (	IMPDH) (II	MPD)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16992775_f2_22	1374	3294	61	186 85	0.00086
Protein name				Locus Name	Acc#
gamma-carboxymuco	onolactone	e decarbo	xylase	pir:B69129	B69129
Description				•	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20353465_f2_21	1375	3295	167	504	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20734687_c2_78	1376	3296	297	894 642	8.2e-63
Protein name				Locus Name	Acc#
				sp:YAAJ_HAEIN	P44555
Description					
HYPOTHETICAL PRO	TEIN HIO1	83			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
21672011_c1_54	1377	3297	61	186 55	0.0095
Protein name				Locus Name sp:YY10_METJA	<u>Acc#</u> Q60309
Description				<u> </u>	<u></u>
HYPOTHETICAL PRO	TEIN MJEC	S10	<u> </u>		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
23439077_f1_5	1378	3298	177	534	103	0.003	5
Protein name ORF MSV035 hypothe	tical p	rotein	, , , , , , , , , , , , , , , , , , ,		s Name 063866		<u>Acc#</u> AF063866
Description							
Melanoplus sanguin	ipes en	comopoxvi	rus, compl	ete geno	me.		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
2347156_f1_8	1379	3299	1105	3318	1839	2.0e-	286
Protein name  isoleucinetRNA I synthetase	igase,::	soleucyl-	-tRNA	Locu:	s Name YECIT		Acc# B64723:S40 549:A94277
<u>Description</u>	NW P	7 7 TD	<u>NT</u>	AA	Caono	Dwaha	:A91325:A9
ORF Name	NTID	AAID	Length	Length	Score		bility
23652183_c1_56	1380	3300	777	2334	3955	0.0	
Protein name Outer membrane pro	tein Cor	n R			s Name		Acc#
Description				gp:U6	9981		U69981
Moraxella catarrha	lis stra	ain Ol2E o	outer memb	rane pro	tein CopP	gene,	complete
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probal	bility
23865660_c2_77	1381	3301	87	264			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	bility
25506316_f1_14	1382	3302	228	687	554	1.7e-	53
Protein name					s Name HA ECOLI		Acc#
Description					_		P24253:P76 771
HYPOTHETICAL GTP-B	INDING	PROTEIN II	N POLA-HEM	N INTERG	ENIC REGI	ON	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	Y
2584717_£3_43	1383	3303	87	264	127	3.1e-08	
Protein name	olactone	e decarbox	ylase	Locu:	s Name	Acc B691	
Description				ــــــــــــــــــــــــــــــــــــــ			
<u>Debel1pe1011</u>							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	Y
25942137_f2_29	1384	3304	185	558	296	3.8e-26	
Protein name				Locus	s Name	Acc	<u>#</u>
				sp:FK	BX_PSEFL	P218	63
Description							
(EC 5.2.1.8) (PPIA	SE) (RO	TAMASE)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabilit	Y
26364431_c1_49	1385	3305	117	354	300	1.4e-26	
Protein name				Locus	s Name	Acc	#
				pir:F	EKRV		
Description						S7216	7:S78 00210
							OULIU
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	Ľ
32056506_c3_81	1386	3306	401	1206	1486	3.0e-152	
Protein name				Locus	Name	Acc	<u>#</u>
				sp:IM	DH_ACICA	P310	02
<u>Description</u>							
DEHYDROGENASE) (IM	PDH) (II	MPD)					
ORF Name	NTID	AAID	NT	<u>AA</u>	Score	Probability	,
			Length	Length			<u> </u>
32477250_c1_65	1387	3307	443	1332	1422	1.8e-145	
Protein name					Name	Acc	_
				sp:YC	DG_ECOLI	P758	92
Description							
HYPOTHETICAL 48.1	KD PROTI	EIN IN WRE	A-PUTA IN	TERGENIC	REGION		

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sc	ore	Probability
34665952_f2_28	1388	3308	177	534	371	4.3e-34
Protein name				Locus N sp:LSPA_		Acc# P17942
Description						
PEPTIDASE) (SIGNA	L PEPTID	ASE II) (S	SPASE II)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sc	ore	Probability
4775762_f1_15	1389	3309	252	759	93	1.3e-57
Protein name				Locus No		Acc# P45528
Description						
HYPOTHETICAL 31.3	KD PROT	EIN IN AGA	AI-MTR INT	ERGENIC REG	ION (F	286)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
5910313_f2_30	1390	3310	392	1179 8	95	1.3e-89
Protein name				Locus Na	ame	Acc#
homoserine O-acet	yltransf	erase		gp:LMMET	YX	Y10744
Description						
L.meyeri metY and	l metX ge	nes.	<u></u> ,			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	Probability
5976592_f2_41	1391	3311	152	459 2	76	5.0e-24
Protein name				Locus Na	ame	Acc#
LporfX				gp:LPU630	641	U63641
Description			<del></del>	J [		
				J [		
Legionella pneumo	phila rpo	oD operon	LporfX, L		_prpoDg	genes, complete
Legionella pneumo cds.  ORF Name	NTID	AAID	NT Length	AA Sco		genes, complete
Legionella pneumo			NT	odnaG, and I		
Legionella pneumo cds.  ORF Name	NTID	AAID	NT Length	AA Sco	ore	
Legionella pneumo cds.  ORF Name  818765_f1_7	NTID	AAID	NT Length	AA Length	ore	Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
9765832_£2_38	1393	3313	458	1377	1102	1.5e-111	
Protein name homoserine dehydrog	enase			Locus gp:L7	s <u>Name</u> 8665	<u>Acc#</u> L78665	
Description							
Methylobacillus fla protein (orf-1), hom thymidylate sythase	noserine	e dehydrog	enase (ho	om), andtl			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
9773436_f2_31	1394	3314	215	648	117	0.00011	
Protein name				Locus	s Name	Acc#	
probable 24-sterol	C-methy	ltransfera	ase,	pir:T	03845	T03845	
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
10423125_c2_44	1395	3315	124	375			
Protein name				Locus	s Name	Acc#	
Description							
NO-HIT		<u> </u>					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
1069202_t2_13	1396	3316	65	198			
Protein name				Locus	s Name	Acc#	
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
12933427_f2_5	1397	3317	131	396	239	4.1e-20	
Protein name				-	Name SC_ECOLI	<u>Acc#</u> P10446	
Description							
SUCCINATE DEHYDROGENASE CYTOCHROME B-556 SUBUNIT							

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20330461_f3_19	1398	3318	229	690	714	1.9e-70
Protein name fumarate reductase	flavopr	otein sub	ınit		s Name 015757	Acc# AB015757
Description						
Rhodoferax fermenta	ns gene	es for fum	arate red	uctase s	ubunits,	complete cds.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
214128_f2_9	1399	3319	763	2292	2566	1.1e-266
Protein name					s Name O1_AZOVI	Acc# P20707
Description						
KETOGLUTARATE DEHYL	ROGENAS	SE)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21501557_f3_27	1400	3320	80	243		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT			•			
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	AA Length	Score	Probability
21510931_f2_6	1401	3321	381	1146	1445	6.6e-148
Protein name				_	s Name	Acc#
fumarate reductase	flavopr	rotein sub	unit	gp:AB	015757	AB015757
Description						
Rhodoferax fermenta	ns gene	es for fum	arate red	luctase s	ubunits,c	complete cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23469010_f3_25	1402	3322	62	189		
Protein name	•			Locu	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23855067_c3_53	1403	3323	183	552	76	0.018
Protein name putative adhesin N	IAA1				s Name 154922	Acc# AF154922
Description						<u>-</u>
Mycoplasma arthricods.	idis st	rain 158 p	outative a	dhesin M	AA1 (maa1	l)gene, complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u>	Score	Probability
24241463_f2_10	1404	3324	68	207	131	2.6e-07
Protein name					s Name O1_HAEIN	Acc# P45303
Description						
KETOGLUTARATE DEH	DROGENA	SE)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24251441_f1_4	1405	3325	375	1128	129	5.8e-05
Protein name				Locus	s Name	Acc#
heme receptor				gp:VI	BHUTA	L27149
Description						
Vibrio cholerae he	eme rece	ptor (hut/	A) gene, c	omplete o	cds.	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24427262_f2_8	1406	3326	123	372	234	2.6e-18
Protein name					s Name	Acc#
alpha-ketoglutarat	e dehyd	rogenase		gp:AF	068740	AF068740
Description						
Pseudomonas putida andalpha-ketogluta						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26367192_f1_2	1407	3327	488	1467	1401	3.0e-143
Protein name				Locus	s Name	Acc#
dihydrolipoamide d	lehydrog	enase		gp:PS	ELPDA	M28356
Description						
P.fluorescens dih	drolipo	amide dehy	/drogenase	(lpd) ge	ene, comp	oletecds.

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
26377042_f2_7	1408	3328	192	579	790	1.7e-	78
Protein name succinate dehydroge sulphur	nase pu	tative iro	on	Locus gp:SP	s Name SDH		<u>Acc#</u> Y13760
Description				_			
Shewanella frigidim	arina N	CIMB400 s	dhA, sdhB	, sdhC, a	sdhD and	sucAge	enes.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
31439375_f1_1	1409	3329	135	408	275	6.3e-	24
Protein name					s Name SD_ECOLI		Acc# P10445
Description						_	
SUCCINATE DEHYDROGE	NASE HY	DROPHOBIC	MEMBRANE	ANCHOR	PROTEIN		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score .	Proba	bility
4064425_f3_26	1410	3330	599	1800	123	8.1e-	05
Protein name					s Name XA_SALTY		<u>Acc#</u> Q56145
Description							
FERRIOXAMINE B RECE	PTOR PR	ECURSOR (	FRAGMENT)			·	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
584625_f2_11	1411	3331	420	1263	1194	2.6e-	121
Protein name				Locus	s Name		Acc#
dihydrolipoamide S-succinyltransferas dehydrogenase comple		-		pir:S	07779		S07779:S63 511
Description							

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
9928130_c1_34	1412	3332	141	426	116	8.6e-07
Protein name				Locu	s Name	Acc#
microfilarial she	eath prote	ein SHP3 p	recursor	gp:AF	030944	
Description						AF030944:U 43510
Brugia malayi mi sheath protein SH			-		_	idmicrofilarial
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
12619081_c3_114	1413	3333	147	444	205	1.7e-16
Protein name					s <u>Name</u> AN_ECOLI	Acc#
Description						P45808:P77 478
HYPOTHETICAL 14.	B KD PROTI	EIN IN PR	IC-APT INT	ERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12897562_c1_73	1414	3334	78	237		
Protein name				Locus	s_Name	Acc#
Description						
NO-HIT	<del></del>					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
1359776_c2_91	1415	3335	67	204		
Protein name				Locus	3 Name	Acc#
Description						
NO-HIT		<u> </u>			·	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14064028_c2_105	1416	3336	269	810	182	4.5e-14
Protein name					Name AB_ECOLI	Acc# P43337
Description					_	
HYPOTHETICAL 21.4	4 KD PROTI	EIN IN PAR	BB-SDAA IN	TERGENIC	REGION	· · · · · · · · · · · · · · · · · · ·

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
14657782_c2_104	1417	3337	173	522 333	4.5e-30
Protein name				Locus Name sp:BID2_HAEIN	Acc# P45248
Description					
2) (DTB SYNTHETASE	2) (DT	BS 2)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14719437_f1_22	1418	3338	63	192	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14882713_c3_116	1419	3339	287	864 289	8.1e-30
Protein name				Locus Name sp:BIOC_HAEIN	Acc# P45249
Description					
PUTATIVE BIOTIN SY	NTHESIS	PROTEIN E	BIOC		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16464750_c2_86	1420	3340	326	981 123	6.0e-05
Protein name				Locus Name sp:ZIPA_ECOLI	Acc# P77173
Description				<u> </u>	
CELL DIVISION PROT	EIN ZIP	Α			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16532256_f1_1	1421	3341	80	243 95	0.0015
Protein name	Licoso			Locus Name	Acc#
ubiquitin protein	TIGase			pir:T39585	T39585
Description					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
19572130_£3_58	1422	3342	310	933 1010	8.2e-102
Protein name				Locus Name sp:CYSM_ECOLI	Acc# P16703
Description					<del></del>
(O-ACETYLSERINE	(THIOL) -L	YASE B) (C	SASE B)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19734630_f2_40	1423	3343	533	1602 400	3.1e-57
Protein name				Locus Name	Acc# P44643
Description					
HYPOTHETICAL RNA	METHYLTR	ANSFERASE	HI0333,		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20507762_f1_13	1424	3344	290	873 548	7.5e-53
Protein name				Locus Name sp:DPSD_ECOLI	Acc# P10740
Description					
PHOSPHATIDYLSERI	NE DECARBO	OXYLASE PR	ROENZYME,		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20839062_c2_92	1425	3345	444	1335 482	1.3e-48
Protein name				Locus Name sp:DEAD_HAEIN	Acc# P44586
Description					
ATP-DEPENDENT RN	A HELICASI	E DEAD HOM	OLOG		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
22144026_f1_26	1426	3346	284	855 443	4.3e-41
Protein name				Locus Name sp:RELA_HAEIN	Acc# P44644
Description	,				
(PPGPP SYNTHETAS)	E I)				

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
22147806_f3_47	1427	3347	421	1266 866	1.5e-86
Protein name	_			Locus Name	Acc# P24196
Description					
(O386)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
22691300_c2_99	1428	3348	612	1839 688	7.1e-72
Protein name				Locus Name	Acc#
sensor kinase rtpA				gp:AB002529	AB002529
Description					
Pseudomonas tolaasi	i gene	for senso	r kinase	rtpA, complete c	ds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
22890917_f2_33	1429	3349	258	777 196	1.5e-15
Protein name  Description				Locus Name	Acc# P52085
HYPOTHETICAL 24.5 F	D PROT	EIN IN PHP	B-HOLA IN	TERGENIC REGION	(ORFUU)
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23478458_c2_103	1430	3350	441	1326	1.7e-135
Protein name				Locus Name	' Acc#
BioA				gp:AF191556	AF191556
Description					
Xenorhabdus nemator (bioA) genes, comple					(var1) and BioA
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24097812_c1_83	1431	3351	208	627 462	9.7e-44
Protein name				Locus Name	Acc# P44409
Description					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24225088_f2_39	1432	3352	61	186		
Protein name	•			Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25587827_c1_79	1433	3353	400	1203	888	7.0e-89
Protein name					s Name OF_HAEIN	Acc# P44422
Description						
LIGASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
26192160_±3_64	1434	3354	532	1599	894	1.6e-89
Protein name					s Name	<u>Acc#</u> P11585
Description				L		
Description (PPGPP SYNTHETASE	: 1)					
	: 1) NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
(PPGPP SYNTHETASE	·	<u>AAID</u>		<u>AA</u>		Probability 4.7e-35
ORF Name	NTID		Length	AA Length 636	Score  380  Name	
ORF Name 29298385_f1_8	NTID		Length	AA Length	Score  380  Name	4.7e-35
ORF Name  29298385_f1_8  Protein name	NTID 1435	3355	Length 211	AA Length 636 Locus gp: U9	Score 380 s Name 0439	4.7e-35 <u>Acc#</u> U90439:AE0 02093
ORF Name  29298385_f1_8  Protein name  Description  Arabidopsis thali	NTID 1435	3355	Length 211	AA Length 636 Locus gp: U9	Score 380 s Name 0439	4.7e-35 <u>Acc#</u> U90439:AE0 02093
ORF_Name  29298385_f1_8  Protein name  Description  Arabidopsis thalisequence.	NTID 1435 ana chroi	3355	Length 211 section 2	AA Length 636 Locus gp:U9	Score  380 S Name 0439	4.7e-35  Acc#  U90439:AE0 02093  complete
ORF Name  29298385_f1_8  Protein name  Description  Arabidopsis thalisequence.  ORF Name	NTID  1435  ana chron  NTID  1436	mosome II  AAID	Length 211 section 2 NT Length	AA Length  AA Locus  AA Length  Locus  Locus  Locus	Score  380 S Name 0439 S of theo	4.7e-35  Acc#  U90439:AE0 02093  complete  Probability
ORF Name  29298385_f1_8  Protein name  Description  Arabidopsis thalisequence.  ORF Name  33708181_c1_75  Protein name	NTID  1435  ana chron  NTID  1436	mosome II  AAID	section 2	AA Length  AA Locus  AA Length  Locus  Locus  Locus	Score  380 S Name 0439 S of theo Score 391 S Name	Acc# U90439:AE0 02093 complete Probability 3.2e-36 Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33728258_c3_117	1437	3357	78	237	85	0.0019
Protein name  Description					s Name D2_HAEIN	Acc# P45248
		RG 2)				
2) (DTB SYNTHETAS	E 2) (DT.	BS 2)		<u>.</u>		
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
35173953_£1_7	1438	3358	152	459	237	6.8e-20
Protein name					s Name EB_ECOLI	Acc#
<u>Description</u>				<u></u>		P05848:P77
HYPOTHETICAL 11.6	KD PROT	EIN IN MRI	A-PHPB IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35183451_f2_38	1439	3359	261	786	259	3.1e-22
Protein name hypothetical prote	ein jhp00	528		Locus	3 <u>Name</u> 71907	Acc# B71907
Description						
ORF Name [4147637_c3_120	NTID	AAID	NT Length	AA Length [2973	Score [3281]	Probability
		3300	390			0.0
Protein name	·				RA_ECOLI	Acc#
Description						P07671:P76 788
EXCINUCLEASE ABC.	SUBUNIT	A				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4199006_f3_56	1441	3361	69	210	52	0.022
Protein name				Locus	Name	Acc#
NADH dehydrogenase	subunit	4		gp:AF	026170	AF026170
Description						
Teius teyou NADH of tRNA-His, tRNA-Ser for mitochondrial p	, and tRI	NA-Leu gen		-	_	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability		
4328431_f3_65	1442	3362	305	918	542	3.2e-52		
Protein name	•				s Name G_NEIME	Acc# P55044		
Description								
GLYCOSYLASE)								
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability		
4867812_c3_118	1443	3363	154	465	318	1.8e-28		
Protein name					s Name HZ_ECOLI	Acc# P32147		
Description								
HYPOTHETICAL 15.9 KD PROTEIN IN RBN-FDHE INTERGENIC REGION (0145)								
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability		
892177 <u></u> c1_70	1444	3364	169	510	331	7.4e-30		
Protein name				Locus gp:D8	s Name	Acc# D83386		
Description				-				
Shewanella violacea	rhIE,	cydD, cyd	C and put	A genes,	partial	andcomplete		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability		
16847336_f3_5	1445	3365	177	534	633	7.3e-62		
Protein name				_	s Name	Acc#		
DNA-directed RNA po.	lymeras	e alpha cl	nain	gp:AF	047025	AF047025		
Description								
Pseudomonas aerugin DNA-directed RNA pol protein L17 (rplQ), bacterioferritin (bf	ymerase and cat	alpha cha alase iso	ain (rpoA zyme A (k	), ribos	omallarge	subunit		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16976442_c3_13	1446	3366	239	720	359	8.0e-33
Protein name					s Name CM_ECOLI	Acc#
Description  HYPOTHETICAL 21.1	VD DROTT		ADD MEDA TA	TPEDCENTA	DECTON	P76938:P76 497
HYPOTHETICAL 21.1	KD PROI	EIN IN F.	-		REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24226655_f2_3	1447	3367	165	498	499	1.2e-47
Protein name				Locu	s Name	Acc#
DNA-directed RNA	polymera	se alpha	chain	gp:AF	047025	AF047025
Description						
Pseudomonas aerug DNA-directed RNA p protein L17 (rplQ) bacterioferritin (	olymeras , and ca	e alpha talase i	chain (rpoA sozyme A (k	), ribos	omallarge	e subunit
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
24317501_f2_4	1448	3368	83	252	354	2.7e-32
Protein name					s Name 17_PSEAE	<u>Acc#</u> 052761
Description						
50S RIBOSOMAL PRO	TEIN L17					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3001693_f2_2	1449	3369	217	654	683	3.7e-67
Protein name				Locus	s Name	Acc#
ribosomal protein	S4			pir:A	64095	A64095
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4867143_c1_9	1450	3370	191	576	314	4.7e-28
Protein name				Locus	s Name	Acc#
probable translat	ion facto	or yciO		pir:F	64874	F64874
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
6033377_c3_14	1451	3371	94	285	84	0.035
Protein name				Locu	s Name	Acc#
hypothetical protei	n C34F6	.9		pir:T	19736	T19736
Description						
ORF Name	NTID .	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10437517_c1_70	1452	3372	62	189		
Protein name		•		Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
11808576_c2_83	1453	3373	72	219		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1359677_£2_17	1454	3374	373	1122	969	1.8e-97
Protein name					s Name	Acc#
uroporphyrinogen de	carboxy	lase		gp:EC	OUW89 	000006
Description						
E. coli chromosomal	region	from 89.	2 to 92.8	minutes	•	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14898317_c3_94	1455	3375	591	1776	1769	3.1e-182
Protein name					s Name	Acc# P21889
Description				25.31		
(ASPRS)						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16522206_c1_56	1456	3376	207	624		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
16614042_c3_107	1457	3377	125	378	142	7.9e-10
Protein name					s Name	Acc#
hypothetical protei	n siri	903		pir:S	77514	S77514
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
175817_f3_32	1458	3378	442	1329	1020	7.2e-103
Protein name				Locu	s Name	Acc#
glyceraldehyde-3-ph	osphate	e dehydrog	enase	gp:AF	058302	AF058302
Description						
Streptomyces roseof sequence.	ulvus	frenolicin	biosynth	etic gen	e cluster	c, complete
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
20984532_c1_68	1459	3379	60	183		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2142151_f3_38	1460	3380	252	759	421	2.1e-39
Protein name				Locus	s Name	Acc#
anion transport ABC homolog ytlC	trans	porter (AT	P-bindi)	pir:C	69995	C69995
Description				-		

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ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23437558_f2_24	1461	3381	348	1047	946	5.0e-95
Protein name  3-phosphoserine a  Description	minotrans	sterase		_	s Name 038578	Acc# AF038578:M 73971:M355 45
Pseudomonas stutz cds;3-phosphoserin dehydratase (aroQp and cyclohexadieny and5-enolpyruvylsh	e aminot: /pheA), ldehydro	ransferase imidazole genase (ty	e (serC), acetolpho rAc) gene	chorisma sphate a es, compl	temutase, minotrans ete cds;	sferase (hisHb),
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
23526888_f1_7  Protein name	1462	3382	65	Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23642875_£3_39	1463	3383	255	768		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24103137_f2_16	1464	3384	409	1230	1079	4.0e-109
Protein name					s Name BZ_HAEIN	Acc# P44915
Description						
HYPOTHETICAL 43.4	KD GTP-F	BINDING PR	OTEIN HIO	877		

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sc	ore	Probability			
24272135_c3_103	1465	3385	174	525	295	4.8e-26			
Protein name				Locus N	ame	Acc#			
Lrp-family transcr	iptiona.	l regulato	rs	gp:D8901	.5	D89015			
Description									
Pseudomonas putida genes for MdeR, MdeA and MdeB, complete cds.									
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sc	ore	Probability			
24410038_f2_19	1466	3386	443	1332	743	1.6e-73			
Protein name				Locus N	ame	Acc#			
proteinase DO				pir:H719	36	H71936			
Description									
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sc	ore	Probability			
25662512_c3_98	1467	3387	304	915	532	3.7e-51			
Protein name				Locus N	ame	Acc#			
				sp:YJJP_	ECOLI	P39402 .			
Description									
HYPOTHETICAL 30.5	KD PROT	EIN IN DNA	T-BGLJ IN	TERGENIC RE	GION (	F277)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	ore	Probability			
25665963_f1_5	1468	3388	264	795 4	144	7.8e-42			
Protein name				Locus Na	ame	Acc#			
				sp:GLO2_	ECOLI	Q47677			
Description									
II) (GLX II)									
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	ore	Probability			
2757750_f3_47	1469	3389	72	219 7	73	0.016			
Protein name				Locus N	ame	Acc#			
				gp:AB021	078	AB021078			
Description						<u>-</u>			
plasmid Collb-P9 D	NA, com	plete sequ	ence.						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31412958_f2_23	1470	3390	250	753		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
33394002_f2_30	1471	3391	507	1524	79	0.036
Protein name				Locu	s Name	Acc#
cytochrome-c oxida	se, cha	in I RP405	<u> </u>	pir:D	71698	D71698
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35181680_c3_95	1472	3392	356	1071	267	4.5e-23
Protein name				Locu	s Name	Acc#
				gp:PF	Y14568	Y14568
Description						
Pseudomonas fluore	scens t	ag gene ar	nd partial	glyQ, h	trB genes	· ,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4009750_c1_69	1473	3393	221	666	279	2.4e-24
Protein name				Locu	s Name	Acc#
hypothetical prote	in			pir:S	76551	S76551
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4165952_f3_34	1474	3394	82	249		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
4328443_c1_74	1475	3395	176	531	193	3.1e-	15
Protein name hypothetical protei	n			Locus pir:G	Name 75479		<u>Acc#</u> G75479
Description							
ORF Name [4423193_c2_79	<u>NTID</u>	<u>AAID</u>	NT Length	AA Length 258	Score	Probal	oility
Protein name	L	J []			Name  GD_ARCFU		Acc# 030156
Description				<u> </u>			
ACETYLORNITHINE AMI	NOTRANS	FERASE, (	ACOAT)	<del></del> .			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probal	oility
4864077_c2_78	1477	3397	63	192	149	1.4e-	10
Protein name				Locus	Name		Acc#
unknown				gp:AF	062531		AF062531
Description						<u> </u>	
Pseudomonas putida unknown genes.	GB-1 si	gnal pept	idase (pi	ID) gene,	partial	. cds;a	nd
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
4878407_f1_6	1478	3398	589	1770	1355	2.3e-1	.38
Protein name					Name J1 YEAST		<u>Acc#</u> P06208
<u>Description</u>				Sp. 1150			100200
SYNTHASE) (ALPHA-IP	M SYNTH	ETASE)					
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probab	oility
5085963_f1_11	1479	3399	243	732	124	3.2e-1	.3
Protein name					Name FN BACSU		Acc# P96692
Description							, , , , <u> </u>
PUTATIVE NAD (P) H NI	TROREDU	CTASE YDFI	Ν,				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5115943_f2_21	1480	3400	330	993	585	9.0e-57
Protein name hypothetical protei	n TM048	34		Locu pir:C	s <u>Name</u> 72369	Acc# C72369
Description				<u> </u>	-	•
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5266588_£2_29	1481	3401	878	2637	2754	1.3e-286
Protein name					s Name	Acc#
UspA1				gp:AF	113606	AF113606
Description						
Moraxella catarrha	lis stra	ain ATCC25	238 UspA1	(uspA1)	gene, co	ompletecds.
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
802137_f3_37	1482	3402	261	786	459	2.0e-43
ABC transporter, pe family	ermease	protein, o	cysTW .	Locus pir:D	72369	<u>Acc#</u> D72369
Description						
ORF Name [894387_c2_80	NTID 1483	<u>AAID</u>	NT Length	AA Length	Score 313	Probability 6.0e-28
Protein name					S Name  UP_HAEIN	Acc# P44520
Description						
HYPOTHETICAL PROTEI	IN HIO10	)8				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
976558_£2_18	1484	3404	61	186		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10740682_c2_12	1485	3405	297	894	678	1.3e-66
Protein name probable acyl-CoA d	ehydrog	genase		Locu:	8 Name 75282	<u>Acc#</u> B75282
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16829202_f3_8	1486	3406	251	753	185	2.2e-14
Protein name					s Name	Acc#
<u>Description</u>				sp:PA	BC_ECOLI	P28305
4-AMINO-4-DEOXYCHOR	ISMATE	LYASE, (A	DC LYASE)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25421887_f1_2	1487	3407	188	567		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT		•				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34160918_f3_7	1488	3408	64	195		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6365631_c3_13	1489	3409	275	828	475	4.1e-45
Protein name				_	s Name	Acc#
shikimate dehydroge	nase			] ab:Nb	U82846	U82846
Description						
Neisseria pharyngis cds.	var. i	iava snik	imate deh	yarogenas 	se (aroE)	gene, complete

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
12156514_c1_16	1490	3410	162	489 336	2.2e-30
Protein name				Locus Name sp:RSTA_ECOLI	Acc# P52108
Description					
TRANSCRIPTIONAL RE	GULATOR	Y PROTEIN	RSTA		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
15625078_c1_19	1491	3411	179	537 444	7.8e-42
Protein name				Locus Name sp:TRMD_SERMA	Acc# P36244
Description				<u> </u>	<del></del>
METHYLTRANSFERASE)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23468928_c1_18	1492	3412	191	576 295	4.8e-26
Protein name				Locus Name sp:RIMM_HAEIN	Acc# P44568
Description					
16S RRNA PROCESSIN	G PROTE	IN RIMM			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23859377_f1_4	1493	3413	502	1509 525	2.0e-50
Protein name				Locus Name	Acc#
EnvZ protein				gp:YEOMPR	Y08950
Description					
Y.enterocolitica o	mpR and	envZ gene	es.		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3961587_c2_21	1494	3414	86	261 279	2.4e-24
Protein name				Locus Name sp:RS16_HAEIN	Acc# P44382
Description					<u>-</u>
30S RIBOSOMAL PROT	EIN S16	<u> </u>			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
964692_c3_22	1495	3415	598	1797	442	1.3e-41
Protein name					s Name TB_ECOLI	Acc# P18392
Description						
SENSOR PROTEIN RST	В,					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10676257_£2_68	1496	3416	393	1182	1212	3.2e-123
Protein name				· · · · · · · · · · · · · · · · · · ·	s Name R9_HAEIN	Acc# P43852
Description				<u> </u>	<del>-</del>	
ORF Name 10736257_f3_80	NTID	<u>AAID</u>	NT Length	AA Length	Score	Probability
Protein name	] [147]	] [3+1/]	03			7 4
Description				Locus	s Name	Acc#
NO-HIT						
			NT	AA	_	
ORF Name	NTID	AAID	Length	Length	Score	Probability
12556337_f1_31	1498	3418	128	387	477	2.5e-45
Protein name		•			s Name	Acc#
Description				sp:PU	R9_ECOLI	P15639
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1272283_f2_60	1499	3419	183	552	244	7.9e-20
Protein name				Locus	s Name	Acc#
Description				sp:AA	RF_ECOLI	P27854:P27 855:P76764 :P27853
UBIQUINONE BIOSYNT	HESIS P	ROTEIN AAR	F			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
131700_c1_108	1500	3420	255	768	215	1.4e-16
Protein name  putative peptidy	-prolyl (	cis-trans	isomerase		Name	<u>Acc#</u> AJ002316
Description	-					<del></del>
Acinetobacter sp	. ADP1 all	kR & alkM	genes, Ok	RF1 & ORF4	1.	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
13876562_c1_128	1501	3421	75	228	73	0.016
Protein name					Name	Acc#
immunoglobulin ka	appa light	chain v	ariabie	gp:AF	131156	AF131156
Description						
Mus musculus immu	unoglobul:	in kappa	light chai	n variab.	le region	n gene, partial
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13947127_c3_217	1502	3422	584	1755	1216	1.6e-163
Protein name					Name	Acc# P43831
Description						,
(GLNRS)						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
14852035_c1_129	1503	3423	85	258	70	0.033
Protein name					Name	Acc#
tat protein				] ab:HI	VU86775	U86775
Description						
protein (rev), vpu complete cds and n	ı protein	(vpu), a	nd envelop	eglycopro		•
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15663417_f2_42	1504	3424	79	240		
Protein name				Locus	Name	Acc#
Description						
NO-HIT	<u> </u>		· · · · · · · · · · · · · · · · · · ·	, <del>12</del> ,,		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16583425_c1_131	1505	3425	326	981	535	1.8e-51
Protein name				Locus	s Name	Acc#
yfjB protein				pir:B	55040	B65040
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19632665_c2_160	1506	3426	696	2091	633	2.3e-79
Protein name					Name PA_ENTHR	Acc#
Description						P32113:Q47 841
COPPER/POTASSIUM-TR	ANSPOR'	ING ATPAS	E A,			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19706557_c3_193	1507	3427	216	651	148	4.8e-08
Protein name	1 1		L	Locus	Name	Acc#
Protein name  probable component cbb3-type oxidase	of cati	on transp	ort for	Locus pir:E		Acc# E71813
probable component	of cati	on transp	ort for			
probable component cbb3-type oxidase  Description  ORF Name	NTID	on transp	NT Length	Pir:E'		
probable component cbb3-type oxidase Description			<u>NT</u>	pir:E'	71813	E71813
probable component cbb3-type oxidase  Description  ORF Name	NTID	AAID	NT Length	AA Length	71813	E71813
probable component cbb3-type oxidase  Description  ORF Name  21753552_c3_220	NTID	AAID	NT Length	AA Length	Score	E71813
probable component cbb3-type oxidase  Description  ORF Name  21753552_c3_220  Protein name	NTID	AAID	NT Length	AA Length	Score	E71813
probable component cbb3-type oxidase  Description  ORF Name  21753552_c3_220  Protein name  Description	NTID	AAID	NT Length	AA Length	Score	E71813
probable component cbb3-type oxidase  Description  ORF Name  21753552_c3_220  Protein name  Description  NO-HIT	NTID 1508	AAID 3428	NT Length 168	AA Length  507 Locus	Score Name	E71813  Probability  Acc#
probable component cbb3-type oxidase  Description  ORF Name  21753552_c3_220  Protein name  Description  NO-HIT  ORF Name	NTID 1508 NTID	AAID  AAID	NT Length 168 NT Length	AA Length  Locus  AA Length  369	Score Name	E71813  Probability  Acc#
probable component cbb3-type oxidase  Description  ORF Name  21753552_c3_220  Protein name  Description  NO-HIT  ORF Name  2197962_f1_11	NTID 1508 NTID	AAID  AAID	NT Length 168 NT Length	AA Length  Locus  AA Length  369	Score  Name	Probability  Acc#  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22145253_c2_177	1510	3430	210	633	592	1.6e-57
Protein name					s Name N_HAEIN	Acc# P45340
Description						
OLIGORIBONUCLEASE,						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22272900_£1_5	1511	3431	227	684	263	1.2e-22
Protein name				Locu	s Name	Acc#
hypothetical prote	in			gp:PS	T243354	AJ243354
Description						
Pseudomonas stutze exbD genes.	ri hypl	and comA	genes and	l putativ	e tolQ, e	exbB,tolR and
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22285902_c3_212	1512	3432	229	690	299	1.8e-26
Protein name				Locus	s Name	Acc#
transposase slr206 slr2062	2:prote	in slr2062	:protein	pir:S	74909	S74909
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22710402_c2_154	1513	3433	78	237		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23457632_f3_90	1514	3434	295	888	885	1.5e-88
Protein name				-	s Name	Acc#
Description				sp:UB	IE_ECOLI	P27851
(EC 2.1.1)						<del>-</del> -

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23475002_f1_9	1515	3435	182	549 121	4.8e-06
Protein name				Locus Name	Acc# P40710
Description					
COPPER HOMEOSTASIS	S PROTEI	N CUTF PRE	CURSOR (I	IPOPROTEIN NLPE)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23554676_f1_16	1516	3436	384	1155 692	4.1e-68
<u>Protein name</u>				Locus Name sp:AARF_ECOLI	Acc#
<u>Description</u>					P27854:P27 855:P76764 :P27853
UBIQUINONE BIOSYN	THESIS P	ROTEIN AAR	EF'		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23634656_c3_200	1517	3437	97	294 147	2.3e-10
Protein name				Locus Name sp:YEAC_ECOLI	Acc# P76231
Description					
HYPOTHETICAL 10.3	KD PROTI	EIN IN ANS	A-GAPA IN	TERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24015950_c1_147	1518	3438	201	606 207	1.0e-16
Protein name				Locus Name	Acc#
hypothetical prote	ein			gp:AF157493	AF157493
Description					
Zymomonas mobilis	ZM4 fost	mid clone	42D7, com	plete sequence.	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24259702_f1_10	1519	3439	500	1503 275	2.4e-23
Protein name				Locus Name sp:YF46 ARCFU	Acc# 028726
Description					
HYPOTHETICAL PROTE	EIN AF154	16			

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24303583_f1_30	1520 3440	93	282	169	1.1e-12
Protein name small DNA binding	protein Fis			s <u>Name</u> 040379	Acc# AF040379
Description					
Proteus vulgaris r cds; yhdG homolog g gene, partial cds.					
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24306510_c3_209	1521 3441	224	675	443	1.0e-41
Protein name				s Name T2_PSEFL	Acc# Q53547
Description					
CARBOXYLESTERASE 2	, (ESTERASE II)				
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24613752_c2_168	1522 3442	241	726	813	6.2e-81
Protein name superoxide dismuta	se, (Mn):SodA pro	tein	Locus pir:J	S Name	<u>Acc#</u> JC6542
Description					<b>'</b>
ORF Name 24614125_t2_65	NTID AAID	NT Length 816	AA Length 2451	Score	Probability
Protein name			Locus	s Name	Acc#
penicillin-binding	protein 1B		gp:AF	147449	AF147449
Description					
Pseudomonas aerugi complete cds.	nosa strain PAO1	penicilli	n-binding	g protein	1 1B(ponB) gene,
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24640762_f3_79	1524 3444	388	1167	1412	2.1e-144
Protein name			· · · · · · · · · · · · · · · · · · ·	Name	Acc#
Description					P04384:P30 869
ADENOSYLTRANSFERAS	E) (ADOMET SYNTHE	TASE)			1

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25551385_f1_17	1525	3445	175	528	298	2.3e-26
Protein name				Locu	s Name	Acc#
adenine phosphoribo	_	sferase,:	protein	pir:S	75440	S75440
sll1430:protein sll1	430				-	
Description				-		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25665962_f1_8	1526	3446	115	348	75	0.0099
Protein name				Locu	s Name	Acc#
glutamyl-tRNA (Gln) C	amidot	ransferase	e subunit	pir:D	70484	D70484
Description		··· -		J		
<u>Deberration</u>						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29687660_±3_88	1527	3447	310	933	441	1.6e-41
Protein name				Locu	s Name	Acc#
hypothetical protei	n			pir:S	76006	S76006
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30650250_f1_12	1528	3448	298	897	664	3.8e-65
Protein name					s Name	Acc#
conserved hypotheti	cal pro	tein		pir:A	75256	A75256
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31541442_c3_183	1529	3449	317	954	338	6.5e-30
Protein name putative peptidyl-p	rolyl c	cis-trans	isomerase	_	s Name AJ2316	Acc# AJ002316
Description		· .				<b>_</b>
Acinetobacter sp. A	DP1 all	kR & alkM	genes, OR	F1 & ORF	4.	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
32694687_c2_181	1530	3450	140	423 117	2.0e-06
Protein name				Locus Name sp:YPBB_BACSU	Acc# P50728
Description					
HYPOTHETICAL 40.	KD PROT	EIN IN FER	-RECQ INT	ERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
33213555_c3_216	1531	3451	60	183 106	1.3e-05
Protein name				Locus Name gp:ECU82664	Acc# U82664
Description					
Escherichia coli	minutes	9 to 11 ge	nomic seq	ruence.	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
33245927_c3_213	1532	3452	229	690 606	5.3e-59
Protein name				Locus Name sp:YCFV_ECOLI	Acc# P75957
Description					
HYPOTHETICAL ABC	TRANSPOR	rer atp-bi	NDING PRO	TEIN YCFV	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
33394050_f3_76	1533	3453	269	810 340	8.2e-31
Protein name				Locus Name sp:YBBF_ECOLI	
Description					P43341:P77
HYPOTHETICAL 26.9	KD PROT	EIN IN PUR	E-PPIB IN	TERGENIC REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
3465_£3_89	1534	3454	71	216	
Protein name				Locus Name	Acc#
Description					
NO-HIT	ALL COLOR		· <del>-</del>	<del></del>	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3906561_f1_18	1535	3455	248	747	311	9.7e-28
Protein name				Locus	Name MBLDA	Acc# M80628
Description						
Streptomyces gris	eus trans	sfer RNA-I	Leu (bldA)	gene and	ORF, co	ompletecds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3910693_£2_39	1536	3456	172	519	526	1.6e-50
Protein name				-	Name PB_ECOLI	Acc#
Description						P23869:P78 052
(ROTAMASE B)		_				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3944178_£2_52	1537	3457	328	987		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3953218_c1_125	1538	3458	943	2832	156	1.1e-10
Protein name				Locus	Name	Acc#
PhoC protein				gp:KPN	1250377	AJ250377
Description						
Klebsiella pneumo PhoC protein.	niae part	cial selD	gene for	SelD prot	ein and	phoCgene for
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3991527_f2_67	1539	3459	2142	6429	577	4.0e-51
Protein name				Locus	Name	Acc# U41852
Description				L		
Haemophilus influ	enzae hsi	gene, co	omplete cd	s.	<del>-</del>	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	ore Pro	bability
4322793_£2_57	1540	3460	217	654		
Protein name				Locus Na	ame	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore Pro	bability
4410943_c3_219	1541	3461	91	276	03 1.1	Le-05
Protein name				Locus Na		<u>Acc#</u> Q46825
Description				<u> </u>		
HYPOTHETICAL 10.5	KD PROT	EIN IN FL	DB-BGLA IN	TERGENIC REC	FION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore Pro	bability
4688887_c3_211	1542	3462	452	1359 1	42 7.9	9e-07
Protein name				Locus Na		Acc#
metal transporter	Nramp4			gp:AF202	540	AF202540
Description						
Arabidopsis thali	ana meta	I transpo	rter Nramp	4 mRNA, comp	olete cds	•
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore Pro	bability
4782812_c1_141	1543	3463	147	444 9	5 0.0	011
Protein name				Locus Na		Acc#
hypothetical prot	ein TM102	26		pir:A7230	)3	A72303
Description						
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Length		bability
4798430_c1_151	1544	3464	453	1362 4	47 7.2	le-62
Protein name				Locus Na		Acc#
Description				gp:SC974!		Z38114:Z71 257
S.cerevisiae chro	mosome X	III cosmi	d 9745.			

ORF Name	NTID AAI	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
5125318_c3_206	1545 34	65 356	1071	184	5.0e-21
Protein name				Name AC007168	Acc# AC007168
Description					
Arabidopsis thalia sequence.	na chromosom	e II BAC T26C	19 genomic	sequenc	ce, complete
ORF Name	NTID AAI	D NT Length	<u>AA</u> Length	Score	Probability
5192757_c1_144	1546 34	66 425	1278	734	1.5e-72
<u>Protein name</u>				Name FW_ECOLI	Acc# P75958
Description					
HYPOTHETICAL 45.3	KD PROTEIN I	N MFD-COBB IN	TERGENIC F	REGION	
ORF Name	NTID AAI	Length	AA Length	Score	Probability
5343752_£3_101	1547 34	316	951	630	1.5e-61
<u>Protein name</u>			-	Name	Acc#
Description			sp:PR	MA_ECOLI	P28637:P76 680:P76681
RIBOSOMAL PROTEIN	L11 METHYLTR	ANSFERASE,			
ORF Name	NTID AAI	- <u>Length</u>	AA Length	Score	Probability
7042580_c1_142	1548 34	75	228		
Protein name			Locus	Name	Acc#
Description					
NO-HIT					
ORF Name	NTID AAI	O NT Length	<u>AA</u> Length	Score	Probability
7312717_£3_77	1549 340	75	228	77	0.028
Protein name conserved hypothet	ical protein	262	Locus	Name 9078	<u>Acc#</u> S59078
Description			<b>-</b> L		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
822680_c3_218	1550	3470	417	1254	675	8.8e-82
Protein name glyceraldehyde-3-ph	osphate	dehydroge	enase		s Name CPGKTIMG	<u>Acc#</u> M87647
Description				<del></del> .		
Bacillus megaterium (gap),phosphoglycera (tpi)genes, complete	te kina					nerase
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
9767325_£3_82	1551	3471	147	444	476	3.2e-45
Protein name transposase homolog	Α			_	s Name U95957	Acc# U95957
Description						
Helicobacter pylori and B (tnpB) genes,		_	nce IS606	transpo	sase homo	ologs A(tnpA)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12635413_f3_5	1552	3472	813	2442	1030	4.2e-129
Protein name					s Name	Acc#
Description						P39170:P39 181:P77465
UNKNOWN PROTEIN FRO	M 2D-PA	GE SPOTS I	M62/M63/O	3/09/T35	PRECURSO	DR .
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
31671880_f1_2	1553	3473	185	558	360	6.2e-33
Protein name				_	s <u>Name</u> U79481	Acc# U79481
Description						
Neisseria meningiti UDP-3-0-(R-3-hydroxy partial cds, and3(R) andUDP-N-acetylgluco	myristo -hydrox	ymyristoy	l acyl ca	rrier pr	otein deh	ydrase (fabZ)

	ORF Name	NTID	AAID	<u>NT</u>	<u>AA</u>	Score	Probability
Protein name					Length		
Description   NO-HIT							
NO-HIT					Locus	Name	Acc#
NTID   AAID   NT   Length	Description						
CRF Name   NTID   AAID   Length   Len	NO-HIT						W 1 W 1
Additional   Add	ORF Name	NTID	AAID			Score	Probability
Sp:LPXA_ECOLI	4412963_t2_4	1555	3475			470	1.4e-44
Description	Protein name				Locus	Name	Acc#
Description					sp:LPX	A_ECOLI	
ORF Name         NTID         AAID         NT Length Length Length         Score         Probability           4687640_f2_3         1556         3476         340         1023         667         1.8e-65           Protein name         Locus Name sp:LPXD_HAEIN         Acc#         P43888           Description         (EC 2.3.1)         NTID AAID NT Length Length Length Length         Score Probability           11978127_f1_2         1557         3477         379         1140         811         1.0e-80           Protein name         Locus Name sp:YECP_ECOLI         P76291:007         983           HYPOTHETICAL 37.0 KD PROTEIN IN ASPS-BISZ INTERGENIC REGION         P76291:007         983           HYPOTHETICAL 37.0 KD PROTEIN IN ASPS-BISZ INTERGENIC REGION         Probability         P76291:007           983         14658562_f3_6         1558         3478         309         930         835         2.9e-83           Protein name         Locus Name sp:YEDI_ECOLI         P46125:P76         3322         P46125:P76	Description						
ORF Name         NTID         AAID         NT Length Length Length         Score         Probability           4687640_f2_3         1556         3476         340         1023         667         1.8e-65           Protein name         Locus Name sp:LPXD_HAEIN         Acc#         P43888           Description         (EC 2.3.1)         NTID AAID NT Length Length Length Length         Score Probability           11978127_f1_2         1557         3477         379         1140         811         1.0e-80           Protein name         Locus Name sp:YECP_ECOLI         P76291:007         983           HYPOTHETICAL 37.0 KD PROTEIN IN ASPS-BISZ INTERGENIC REGION         P76291:007         983           HYPOTHETICAL 37.0 KD PROTEIN IN ASPS-BISZ INTERGENIC REGION         Probability         P76291:007           983         14658562_f3_6         1558         3478         309         930         835         2.9e-83           Protein name         Locus Name sp:YEDI_ECOLI         P46125:P76         3322         P46125:P76	[/EC 2 2 1 128\ /M	DD-N-XCE		CAMTNE ACVI	TOANGEEDA	ਵਜ਼।	
NTID   AAID   Length   P43888      Description     Description   Descri	(EC 2.3.1.129) (0	DP-N-ACE	11101000				
Protein name	ORF Name	NTID	AAID			Score	Probability
Description   Sp:LPXD_HAEIN   P43888	4687640_f2_3	1556	3476	340	1023	667	1.8e-65
Description   CEC 2.3.1)   ORF Name   NTID   AAID   Length   P76291:007	Protein name				Locus	Name	Acc#
ORF Name					sp:LPX	D_HAEIN	P43888
ORF Name         NTID         AAID         NT Length         Length         Score         Probability           11978127_f1_2         1557         3477         379         1140         811         1.0e-80           Protein name         Locus Name         Acc#           Sp:YECP_ECOLI         P76291:007           983         P76291:007           HYPOTHETICAL 37.0 KD PROTEIN IN ASPS-BISZ INTERGENIC REGION           ORF Name         NTID         AAID         NT         AAA         Score         Probability           14658562_f3_6         1558         3478         309         930         835         2.9e-83           Protein name         Locus Name         Acc#           Sp:YEDI_ECOLI         P46125:P76           332	Description						
NTID   AAID   Length   Length   Score   Probability	(EC 2.3.1)		<del></del>				
Description	ORF Name	NTID	AAID			Score	Probability
Sp:YECP_ECOLI	11978127_f1_2	1557	3477			811	1.0e-80
Sp:YECP_ECOLI	Protein name				Locus	Name	Acc#
Description   983							
HYPOTHETICAL 37.0 KD PROTEIN IN ASPS-BISZ INTERGENIC REGION	Description					····	
ORF Name         NTID         AAID         NT Length         Length         Score         Probability           14658562_f3_6         1558         3478         309         930         835         2.9e-83           Protein name         Locus Name         Acc#           Sp:YEDI_ECOLI         P46125:P76           332         332							
Name   NTID   AAID   Length   Length	HYPOTHETICAL 37.0	KD PROT	EIN IN A	SPS-BISZ IN		REGION	
Protein name  Locus Name  Sp:YEDI_ECOLI  P46125:P76  332	ORF Name	NTID	AAID			Score	Probability
Description P46125:P76	14658562_f3_6	1558	3478	309		835	2.9e-83
Description P46125:P76	Protein name				Locus	Name	Acc#
Description 332					sp:YED	I_ECOLI	D46105 B56
HYPOTHETICAL 32.2 KD PROTEIN IN DSRB-VSR INTERGENIC REGION	Description						
		KD PROT	EIN IN D	SRB-VSR INT	ERGENIC R	EGION	1

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23714375_f3_8	1559	3479	100	303 70	0.033
Protein name outer membrane pro	tein H.	8 precurso	r	Locus Name	Acc# S04157
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24253427_f3_7	1560	3480	85	258	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24322153_f3_5	1561	3481	257	774 475	4.1e-45
Protein name				Locus Name	Acc#
Description					P43985:P43 986
HYPOTHETICAL PROTE	IN HIO3	19/320			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24804651_c2_16	1562	3482	62	189 171	3.1e-12
Protein name				Locus Name sp:SSP2 PLAYO	<u>Acc#</u> Q01443
Description					~
SPOROZOITE SURFACE	PROTEI	N 2 PRECUR	SOR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
35181956_c1_11	1563	3483	251	756	
Protein name				Locus Name	Acc#
D = = = = 1 = 1 = =					
Description					<del></del>

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3955437_c2_19	1564	3484	73	219	138	2.1e-09
Protein name				Locu	s Name	Acc#
peptide methionine	sulfoxi	de reduct	ase	pir:E	75345	E75345
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5117337_£3_9	1565	3485	379	1140	1148	2.0e-116
Protein name serine-pyruvate ami	notrans	férase		Locu:	s <u>Name</u> 75269	Acc# F75269
Description				J <u>E</u>		
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	AA Length	Score	Probability
1063441_c3_193	1566	3486	442	1329	272	1.4e-21
Protein name			_		s Name	Acc#
hypothetical protei	n 25			pir:T	13514	T13514
Description						
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
10650681_c3_218	1567	3487	102	309	72	1.0e-05
Protein name					s Name	Acc#
unknown				gp:AF	050676	AF050676
Description						
Pseudomonas aerugin genes, complete cds;	_	-	_	d ferric	uptakere	egulator (fur)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
119027_c2_166	1568	3488	95	288		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT		<del>-,</del>				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1227302_c3_221	1569	3489	90	273	85	0.012
Protein name probable fatty-ac	cidCoA l	igase,	adD7	Locus pir:C	s Name 69471	Acc# C69471
Description						
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Length	Score	Probability
12773910_c2_171	1570	3490	116	351		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12973332_f1_18	1571	3491	63	192		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12992125_c2_170	1572	3492	152	459		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13085160_c3_196	1573	3493	236	711		
Protein name				Locus	s_Name	Acc#
Description						
NO-HIT					<del></del>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1371003_£3_87	1574	3494	413	1242	1088	4.5e-110
Protein name				Locus	s Name	Acc#
Na+/H+-exchanging	protein:	Na+/H+ a	antiporter	pir:J	X0360	JX0360
Description				<del></del>		

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14647033_c3_209	1575	3495	192	579 128	1.8e-07
Protein name				Locus Name	Acc#
muramoyl-pentapepti	de carl	poxypeptid	ase	pir:T34747	T34747
Description				·	
ORF Name	NTID	AAID .	<u>NT</u> Length	AA Length Score	Probability
14745253_c3_214	1576	3496	468	1407 596	6.1e-58
Protein name				Locus Name	Acc#
hypothetical protei	n Rv37.	34c		pir:G70797	G70797
Description				_	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
15633253_c1_149	1577	3497	150	453	
Protein name				Locus Name	Acc#
<u>Description</u>					
NO-HIT					
ORF Name	NTID	AAID	NT	AA Score	Probability
16485906 c3_192	1578	3498	Length 484	<u>Length</u> 1455 207	2.3e-14
	] [23,0		101	L	
Protein name				<u>Locus Name</u> sp:VG17_BPMD	Acc# 2 064210
Description				<u> </u>	001210
MAJOR HEAD PROTEIN	GP17	<del></del>			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
16595716_£2_69	1579	3499	60	183	
Protein name				Locus Name	Acc#
Description					
NO-HIT					

	AA Score Probability
16597827_c1_133	618 92 0.022
Protein name putative prohead protease	<u>Locus Name</u> <u>Acc#</u> <u>gp:AF181080</u> AF181080
Description	
Rhodobacter capsulatus putative large termi putative prohead protease genes, complete cogene, partial cds.	
ORF Name NTID AAID NT Length	AA Length Score Probability
19547875_c1_157	375 189 8.2e-15
Protein name	Locus Name Acc#
mono-heme c-type cytochrome ScyA	gp:AF044582 AF044582
Description	
Shewanella putrefaciens NrfG homolog gene, cytochrome ScyA (scyA), cytochrome c maturat maturation protein B (ccmB),cytochrome c matcytochrome c maturationprotein D (ccmD), and E (ccmE)genes, complete cds.	cionprotein A (ccmA), cytochrome c curation protein C (ccmC),
ORF Name NTID AAID NT Length	AA Length Score Probability
19697265_c2_179	198 75 0.020
Protein name	Locus Name Acc#
	sp:YC67_ASTLO P34778
Description	
HYPOTHETICAL 20.1 KD PROTEIN YCF67 (ORF170)	
ORF Name NTID AAID NT Length	AA Length Score Probability
20917082_f3_105	216
Protein name	Locus Name Acc#
Description	
ро-ніт	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	ore Prob	ability
21663410_f1_17	1584	3504	170	513		
Protein name				Locus Na	ıme	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	ore Prob	ability
22351557_c3_191	1585	3505	89	270	0.00	39
Protein name hypothetical protei	n 2068	. 13		Locus Na	<del></del>	Acc#
	11 FZ6B6			pir:T0114		T01147
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	re Prob	ability
22381542_c3_199	1586	3506	257	774 4	3.3e	-43
Protein name				Locus Na		Acc#
minor tail protein	L homol	.og:protei	n gp18	pir:T1310	04	T13104
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	re Proba	ability
23437561_±2_52	1587	3507	690	2073	985 4.0e	-205
Protein name				Locus Na	me	Acc#
				sp:SYM_HA	EIN	P43828
Description						
(METRS)	<del></del>					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	re Proba	ability
23549217_c1_144	1588	3508	192	579 1:	5.4e	- 05
Protein name				Locus Na	me	Acc#
hypothetical protei	n			pir:T1465	1	T14651
Description				_		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	core	Probab	ility
23847257_c2_167	1589	3509	125	378			
Protein name				Locus	Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probab	ility
24316886_f3_115	1590	3510	151	456	356	1.7e-3	2
Protein name				Locus 1			Acc#
Description				sp:YDCQ	ECOLI		P76107
HYPOTHETICAL 16.1	KD PROT	EIN IN TE	HB-ANSP IN	TERGENIC R	EGION		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probab	ility
24415876_f2_48	1591	3511	154	465			
Protein_name				Locus 1	Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length So	core	Probab	ility
24417540_c2_187	1592	3512	209	630	600	2.3e-5	8
Protein name				Locus 1			Acc#
				gp:XCRP	FB		Y09700
Description							
X.campestris rpfB	gene.						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probab	ility
24431265_c2_182	1593	3513	486	1461	1261	2.1e-1	28
Protein name				Locus 1			Acc#
<u>Description</u>				sp:SYC_	ECOLI		P21888
(CYSRS)							

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
24614431_c2_173	1594	3514	169	510	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24631552_c2_181	1595	3515	271	816 723	2.1e-71
Protein name				Locus Name	Acc#
thiamin biosynthe	sis prote	ein thiG		pir:B70487	B70487
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24882676_c2_163	1596	3516	198	597 261	1.9e-22
Protein name				Locus Name	Acc#
				sp:YE18_HAEIN	P44189
Description					
HYPOTHETICAL PROT	EIN HI14	18			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
25397700_c1_148	1597	3517	221	666 388	6.7e-36
Protein name				Locus Name	Acc#
minor tail protei	n gp20			pir:T13106	T13106
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
25493762_c1_161	1598	3518	60	183	
Protein name				Locus Name	Acc#
Description					
NO-HIT					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25584627_c3_216	1599	3519	60	183		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT	V.,	<u></u>	, <del></del>			7.77
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
2581542_c3_213	1600	3520	72	219		
Protein name				Locu	s Name	Acc#
Description				<u></u>		
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26819002_c1_141	1601	3521	90	273	72	0.020
Protein name				Locu	s Name	Acc#
hypothetical prote	in yorB			pir:T	12887	
hypothetical prote Description	in yorB			pir:T	12887	T12887:C69
	in yorB	AAID	<u>NT</u> . Length	pir:T  AA  Length	12887 <u>Score</u>	· ·
Description		<u>AAID</u>		<u>AA</u>		922
Description  ORF Name	NTID		Length	AA Length 993	Score	922 Probability
Description  ORF Name  276927_f2_58  Protein name	NTID		Length	AA Length 993 Locus	Score	922 Probability 0.0016
Description  ORF Name  276927_f2_58	NTID		Length	AA Length 993 Locus	Score [111] s Name	922 Probability 0.0016 Acc#
Description  ORF Name  276927_f2_58  Protein name	NTID		Length	AA Length 993 Locus	Score [111] s Name	922 Probability 0.0016 Acc#
Description  ORF Name  276927_f2_58  Protein name  Description	NTID		Length	AA Length 993 Locus	Score [111] s Name	922 Probability 0.0016 Acc#
Description  ORF Name  276927_f2_58  Protein name  Description  FINQ PROTEIN	NTID 1602	3522	Length 330	AA Length 993 Locus Sp:FI	Score  [111] s Name NQ_ECOLI	922  Probability  0.0016  Acc# P18809
ORF Name  276927_f2_58  Protein name  Description  FINQ PROTEIN  ORF Name	NTID 1602 .	3522 AAID	Length  330  NT Length	AA Length Sp:FI  AA Length  AA Length  339	Score  [111] S Name  NQ_ECOLI  Score  [112] S Name	Probability  O.0016  Acc# P18809  Probability  1.2e-06  Acc#
Description  ORF Name  276927_f2_58  Protein name  Description  FINQ PROTEIN  ORF Name  2792176_c2_180	NTID 1602 .	3522 AAID	Length  330  NT Length	AA Length Sp:FI  AA Length  AA Length  339	Score  [111] S Name  NQ_ECOLI  Score  [112]	Probability  O.0016  Acc# P18809  Probability  1.2e-06

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29337908_f1_37	1604	3524	82	249		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT				11.0		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
31678827_c3_222	1605	3525	244	735	527	1.3e-50
Protein name  long-chain-fatty-a	cid-CoA	ligase			s Name 150669	<u>Acc#</u> AF150669
Description						
Pseudomonas putida cds.	long-ch	nain-fatty	-acid-CoA	ligase	(fadD) ge	ene, complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
32062552_f3_102	1606	3526	61	186	54	0.0065
Protein name					s Name 83_METJA	<u>Acc#</u> Q58096
Description				<u>-</u>		
HYPOTHETICAL PROTE	IN MJ068	33				
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
3207751_f3_103	1607	3527	126	381		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
35187543_c3_217	1608	3528	378	1137	105	0.0058
Protein name AdcB protein	-			Locu gp:SP	s Name ADCA	Acc# Z71552
Description		-	W			
Streptococcus pneu	moniae a	adcRCBA or	eron.			

ORF Name	NTID AAID	NT AA Length Length Scor	e <u>Probability</u>
3937813_c1_158	1609 3529 2	233 702 368	8.9e-34
Protein name		Locus Nam	
Description			
CYTOCHROME C4 PREC	URSOR		
ORF Name	NTID AAID	NT <u>AA</u> Length <u>Length</u> Scor	e Probability
402217_c3_194	1610 3530	486	
Protein name		Locus Nam	e <u>Acc#</u>
Description			
NO-HIT			
ORF Name	NTID AAID	<u>NT AA</u> Length Length Scor	e <u>Probability</u>
4069212_c3_195	1611 3531	357 83	0.017
Protein name	•	Locus Nam sp:Y182 ME	<del></del>
Description		L	
HYPOTHETICAL PROTE	IN MJ0182		
ORF Name	NTID AAID	<u>NT AA</u> Length Length	e Probability
4331563_c2_172	1612 3532	3540 181	1.2e-09
Protein name		Locus Nam	
unknown		gp:AF01137	8 AF011378
Description			
Bacteriophage skl	complete genome.		
ORF Name	NTID AAID	<u>NT AA</u> Length <u>Length</u> Scor	e <u>Probability</u>
4415938_c2_177	1613   3533   1	1627 4884 186	3.4e-198
Protein name tail tip fiber pro	otein gp21	Locus Nam	<del>-</del>

ORF Name	NTID	AAID	Length	<u>AA</u> Length	Score	Probability
4861263_c2_169	1614	3534	121	366		
Protein name		_		Locus	Name	Acc#
Description						
NO-HIT		*	<del></del>	·		<u>,                                      </u>
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4867819_c2_162	1615	3535	196	591	404	1.4e-37
Protein name				Locus	Name	Acc#
hypothetical prot	ein HP133	4		pir:F6	4686	F64686
Description				<del></del>		<u>-</u>
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5130075_c2_186	1616	3536	431	1296	858	1.1e-85
Protein name				Locus	Name	Acc#
Protein name		J [			Name P_HAEIN	Acc# P44953
Protein name  Description	<b>-</b>	J I				
	METABOLIS	SM FLAVOPR	<del> </del>	sp:DFI		
Description	METABOLIS NTID	EM FLAVOPR	<del> </del>	sp:DFI		
Description DNA/PANTOTHENATE			OTEIN HON	sp:DFF OLOG <u>AA</u>	P_HAEIN	P44953
Description  DNA/PANTOTHENATE  ORF Name	NTID	AAID	OTEIN HON  NT  Length	Sp:DFI OLOG  AA Length 276	P_HAEIN	P44953
Description  DNA/PANTOTHENATE  ORF Name  553437_f1_28	NTID	AAID	OTEIN HON  NT  Length	Sp:DFI OLOG  AA Length 276	Score	P44953  Probability
Description  DNA/PANTOTHENATE  ORF Name  553437_f1_28  Protein name	NTID	AAID	OTEIN HON  NT  Length	Sp:DFI OLOG  AA Length 276	Score	P44953  Probability
Description  DNA/PANTOTHENATE  ORF Name  553437_f1_28  Protein name  Description	NTID	AAID	OTEIN HON  NT  Length	Sp:DFI OLOG  AA Length 276  Locus	Score	P44953  Probability
Description  DNA/PANTOTHENATE  ORF Name  553437_f1_28  Protein name  Description  NO-HIT	NTID 1617	<u>AAID</u> 3537	NT Length 91	Sp:DFI OLOG  AA Length 276  Locus	Score	P44953  Probability  Acc#
Description  DNA/PANTOTHENATE  ORF Name  553437_f1_28  Protein name  Description  NO-HIT  ORF Name	NTID 1617 NTID	AAID 3537 AAID	OTEIN HON  NT Length  91  NT Length	Sp:DFI OLOG  AA Length  Locus  AA Length  4A Length	Score Score	Probability  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
682777_c1_145	1619	3539	135	408		
Protein name				Locus	s Name	Acc#
Description		•				
NO-HIT			<del></del>			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
683187_c1_135	1620	3540	71	216		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT					· · ·	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6925452_£3_100	1621	3541	68	207	69	0.042
		_				
Protein name				Locus	Name	Acc#
Protein name hypothetical pro	tein APEO	740		Locus		<u>Acc#</u> E72664
	tein APEO'	740				
hypothetical pro	tein APEO	AAID	<u>NT</u> Length			
hypothetical pro Description			<del></del>	pir:E'	72664	E72664
hypothetical pro Description ORF Name	NTID	AAID	Length	AA Length	72664	E72664
hypothetical pro Description  ORF Name 790807_f1_16	NTID	AAID	Length	AA Length	Score	E72664  Probability
hypothetical pro Description  ORF Name 790807_f1_16 Protein name	NTID	AAID	Length	AA Length	Score	E72664  Probability
hypothetical pro  Description  ORF Name  790807_f1_16  Protein name  Description	NTID	AAID	Length	AA Length	Score	E72664  Probability  Acc#
hypothetical pro  Description  ORF Name  790807_f1_16  Protein name  Description  NO-HIT	NTID 1622	<u>AAID</u> 3542	Length 101 NT	AA Length  306  Locus	Score  Name	E72664  Probability  Acc#
hypothetical pro  Description  ORF Name  790807_f1_16  Protein name  Description  NO-HIT  ORF Name	NTID 1622 NTID	<u>AAID</u> 3542  AAID	Length 101 NT Length	AA Length  Locus  AA Length  198	Score  Name	E72664  Probability  Acc#
hypothetical pro  Description  ORF Name  790807_f1_16  Protein name  Description  NO-HIT  ORF Name  830300_f1_21	NTID 1622 NTID	<u>AAID</u> 3542  AAID	Length 101 NT Length	AA Length  Locus  AA Length  198	Score  Score	Probability  Acc#  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Scor	re <u>Probability</u>
865782_c3_198	1624	3544	750	2253 17	3 6.3e-12
Protein name				Locus Nar	
Description					
Pseudomonas aeru	ginosa ger	nomic DNA	, partial	sequence, str	rain:PAO1.
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Scor	e Probability
14175056_±1_2	1625	3545	67	204 11	6 4.5e-07
Protein name				Locus Nam	_
Description					<del></del>
A.brasilense car	R gene.				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Scor	re Probability
23831527_c3_33	1626	3546	674	2025 61	9 1.7e-74
Protein name protein-disulfide	e reductas	se		Locus Nam	<del></del>
Description				J [=-	
Pseudomonas aeru dehydroquinase (a				uctase (dipZ)	andcatabolic
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Scor	e Probability
26276961_c2_28	1627	3547	405	1218 16	07 4.5e-165
Protein name				Locus Nam	ne Acc#
chloroacetaldehy	de dehydro	ogenase		gp:AF02973	AF029733
Description					
Xanthobacter aut chloroacetaldehyd	_	_			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Scor	e Probability
33581289_c2_24	1628	3548	512	1539 12	11 4.1e-123
Protein name				Locus Nam	<del>-</del>
Description					<del></del>
HYPOTHETICAL SOD	IUM-DEPENI	DENT TRANS	SPORTER HI	0736	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5312692_f3_15	1629	3549	411	1236	1075	1.1e-108
Protein name				Locu	s Name	Acc#
sodium/proton-deper	ndent al	anine cari	rier pr	pir:C	69972	C69972
homolog yrbD					-	
Description				_		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6152307 c2 26	1630	3550	387	1164	1087	5.7e-110
Protein name	J L	لـــــا لــ	LJ	Locus	s Name	Acc#
FIOCEIII IIamo					DB_ECOLI	P11027
Description				L.		
BD-I OXIDASE SUBUN	IT II)					
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
781461_c2_25	1631	3551	480	1443	1563	2.1e-160
Protein name	<b>-</b>	J <u>L</u>		Locu	s Name	Acc#
				sp:CY	DA_AZOVI	Q09049
Description						
CYTOCHROME D UBIQU	INOL OXI	DASE SUBU	NIT I,			
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
125143_c1_36	1632	3552	82	249	137	5.7e-09
Protein name				Locu	s Name	Acc#
probable enoyl-CoA	hydrata	se		pir:G	75557	G75557
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12632255_c3_48	1633	3553	239	720	136	8.4e-06
Protein name				Locu	s Name	Acc#
probable erythrocy	te-bindi	ng protein	n MAEBL	pir:T	09127	T09127
Description						

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability
13064425_f1_6	1634	3554	160	483	569	4.4e-55
Protein name					Name M6_ECOLI	Acc# P36553
Description						
(COPROPORPHYRINOGE	NASE) (C	OPROGEN O	XIDASE)		-	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16692186_£2_20	1635	3555	158	477	166	2.3e-12
Protein name				-	Name CP_ALCSP	Acc# P00138
Description				L		·
CYTOCHROME C'						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
195277_f1_11	1636	3556	405	1218	491	8.2e-47
				L		
Protein name					Name	Acc#
Protein name ORF396 protein				Locus gp:PSI		Acc# Z73914
ORF396 protein	eri ort17	5 gene.				
ORF396 protein Description	eri ort17	5 gene.	NT Length			
ORF396 protein  Description  Pseudomonas stutze				gp:PSI	ONGC	Z73914
ORF396 protein  Description  Pseudomonas stutze  ORF Name	NTID	AAID	Length	AA Length 2133 Locus	Score	Z73914  Probability
ORF396 protein  Description  Pseudomonas stutze  ORF Name  197137_c2_46	NTID	AAID	Length	AA Length 2133 Locus	Score  965  Name	Z73914  Probability  2.5e-156  Acc#
ORF396 protein  Description  Pseudomonas stutze  ORF Name  197137_c2_46  Protein name	NTID 1637	<u>AAID</u> 3557	Length 710	AA Length 2133 Locus sp:DX	Score  965  Name	Z73914  Probability  2.5e-156  Acc#
ORF396 protein  Description  Pseudomonas stutze  ORF Name  197137_c2_46  Protein name  Description	NTID 1637	<u>AAID</u> 3557	Length 710	AA Length 2133 Locus sp:DX	Score  965  Name	Z73914  Probability  2.5e-156  Acc#
ORF396 protein  Description  Pseudomonas stutze  ORF Name  197137_c2_46  Protein name  Description  1-DEOXYXYLULOSE-5-	NTID 1637 PHOSPHAT	AAID 3557 E SYNTHAS	Length 710 E (DXP SY	AA Length 2133 Locus sp:DX	Score  [965] S Name S HAEIN	Z73914  Probability  2.5e-156  Acc# P45205
ORF396 protein  Description  Pseudomonas stutze  ORF Name  197137_c2_46  Protein name  Description  1-DEOXYXYLULOSE-5-  ORF Name	NTID  1637  PHOSPHAT  NTID	AAID  3557  E SYNTHAS  AAID	Length 710  E (DXP SY  NT  Length	AA Length 2133 Locus Sp:DXS NTHASE) AA Length 315	Score  965  Name S_HAEIN	Probability  2.5e-156  Acc# P45205  Probability
ORF396 protein  Description  Pseudomonas stutze  ORF Name  197137_c2_46  Protein name  Description  1-DEOXYXYLULOSE-5- ORF Name  22697263_c1_37	NTID 1637 PHOSPHAT NTID 1638	AAID  E SYNTHAS  AAID  3558	Length 710  E (DXP SY  NT  Length	AA Length 2133 Locus Sp:DXS NTHASE) AA Length 315	Score  965 S Name S HAEIN  Score  87 S Name	Probability  2.5e-156  Acc# P45205  Probability  0.0022

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Proba	bility
24323500_f1_5	1639	3559	171	516	435	7.0e-	41
Protein name					s Name M6_ECOLI		<u>Acc#</u> P36553
Description							
(COPROPORPHYRINOG	ENASE) (	COPROGEN	OXIDASE)				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
30120325_c1_32	1640	3560	103	312			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
33449042_c3_54	1641	3561	126	378	294	6.2e-	26
Protein name				Locus	s Name		Acc#
SuhB				gp:AF	010139		AF010139
Description							
Azotobacter vinel cysE2, iscS, iscU, partial cds.							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
33986312_f2_16	1642	3562	209	630	522	4.3e-	50
Protein name					Name H2_HAEIN		Acc# P44571
Description							
GTP CYCLOHYDROLAS	E II,		<del></del>				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
35441086_c2_43	1643	3563	149	450	96	0.011	
Protein name					Name		Acc#
cell wall-binding	protein	homolog	YVCE	pir:F	70031		F70031
Description							

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
5859703_c1_33	1644	3564	464	1395 705	1.7e-69
Protein name				Locus Name	Acc# J02808
Description					
E.coli folC gene er protein required for					synthetase, and a
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
1046926_c1_177	1645	3565	217	654 325	3.2e-29
Protein name				Locus Name	Acc#
yrp protein:multipl	.e regu.	lator prot	ein	pir:S70842	\$70842
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
10588311_c3_274	1646	3566	401	1206 1495	3.3e-153
Protein name				Locus Name	Acc#
ribonucleoside-diph chain	osphate	reductas	e, beta	pir:C64135	C64135
Description				_	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
10602250_£2_95	1647	3567	132	399 251	2.2e-21
Protein name				Locus Name	Acc#
aluminum tolerance	proteir	1		pir:PC4440	PC4440:PC4
Description					
					514
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	514 Probability
ORF Name 10751006_c1_182	<u>NTID</u>	AAID 3568		— Score	
			Length	Length Score	Probability
10751006_c1_182			Length	Length Score  Locus Name	Probability  5.0e-08  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
11912951_f1_20	1649	3569	107	324		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1297216_c3_289	1650	3570	185	558	125	5.0e-08
Protein name				Locu	s Name	Acc#
colicin V producti	on prote	ein homol	og 	pir:E	70195	E70195
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14275330_t2_68	1651	3571	489	1470	377	9.9e-35
Protein name				Locus	s Name	Acc#
Description				sp:Y4	WB_RHISN	P55680
HYPOTHETICAL ZINC	PROTEAS:	E-LIKE PR	OTEIN Y4WB			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14508500_c2_247	1652	3572	513	1542	1546	1.3e-158
Protein name amidophosphoribosy	Itransf	erase			s Name	Acc#
				pir:X	QEC .	F65003:A92
<u>Description</u>						366:A92367 :S01389:I5
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14900187_f3_134	1653	3573	220	663	361	4.9e-33
Protein name				_	s Name	Acc#
probable 2-hydroxy isomerase b1180	hepta-2	4-diene-	1,7-dioate	pir:A	64864	A64864
Description	<del></del>			J		

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15908263_c1_143	1654 3574	108	327	307	2.6e-27
Protein name RpsA			_	Name 035937	Acc# AF035937
Description					
Pseudomonas aerugir Ihf-Beta, Wzz (wzz), for O-antigen biosyr	, and Wzx (wzx) g	enes, com	plete cds	_	
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16194442_f3_135	1655 3575	458	1377	1285	6.0e-131
Protein name			-	Name	<u>Acc#</u> P26977
Description					
RIBONUCLEOTIDE SYNT	THETASE) (PHOSPHO	RIBOSYLGL	YCINAMIDE	SYNTHE	TASE)
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16828790_f1_43	1656 3576	292	879	398	5.9e-37
Protein name				Name D_HAEIN	Acc# P44710
Description					
HYPOTHETICAL PROTEI	N H10432		***		
ORF Name 19698381_c1_189	NTID <u>AAID</u>	NT Length	AA Length	Score	Probability
Protein name Description			Locus	Name	Acc#
NO-HIT					
		NT	AA		
ORF Name	NTID AAID	Length	Length	Score	Probability
1972931_£2_63	1658 3578	68	207	57	0.023
Protein name unknown			Locus gp:AF1		Acc# AF197128
Description					
Rattus norvegicus u	inknown mRNA.			<del> </del>	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20601558_c1_163	1659	3579	273	822	737	7.0e-73
Protein name					Name U_ECOLI	<u>Acc#</u> Q46845
Description						
HYPOTHETICAL 34.2	KD PROT	EIN IN GSP	-HYBG INT	ERGENIC R	EGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2111556_c1_164	1660	3580	1383	4152	161	1.3e-29
Protein name				Locus sp:EX5	Name C_HAEIN	Acc# P44945
Description				<del></del>		
EXODEOXYRIBONUCLE	ASE V GA	MMA CHAIN,				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
21642556_c3_272	1661	3581	87	264		
Protein name				Locus	Name	Acc#
Description						
NO-HIT			-			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
22751387_c3_271	1662	3582	150	453		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23611527_c3_275	1663	3583	114	345	140 ′	1.3e-09
Protein name				Locus sp:YFA	Name E_HAEIN	Acc# P45154
Description				<u></u>		
HYPOTHETICAL PROT	EIN HI13	09		<del> </del>		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23676035_c3_262	1664	3584	410	1233	177	1.3e-10
Protein name YtfP		-		_	s Name 008220	Acc# AF008220
Description						
Bacillus subtilis	rrnB-dna	aB genomic	region.			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23725387_c2_244	1665	3585	318	957	1046	1.3e-105
Protein name					S Name	Acc# P44870
Description						
CELL DIVISION PRO	TEIN FTS	7				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23728465_c1_161	1666	3586	925	2778	2856	2.0e-297
Protein name				_	s Name	Acc#
pyruvate dehydrog	enase (li	.poamide)		gp:AZ	PDHE	Y15124
Description						
Azotobacter vinel	andii pdh	E gene.				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
23989512_c3_268	1667	3587	393	1182	1066	9.6e-108
Protein name					s Name	Acc#
<u>Description</u>				sp:PH	EA_PSEST	P27603
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24303127_c1_171	1668	3588	407	1224	797	3.1e-79
Protein name carboxynorspermid	ine decar	boxylase		_	s Name BCANSDC	Acc# D31783
Description				T Br.AT		231/63
Vibrio alginolyti DC), complete cds.	cus nspC	gene for	carboxyno	rspermid	inedecarb	oxylase (CANS

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24407750_c3_253	1669	3589	248	747 603	1.1e-58
Protein name				Locus Name	Acc# P43812
Description					
DECARBOXYLASE)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24642711_c2_216	1670	3590	773	2322 960	1.6e-96
Protein name				Locus Name sp:AROA_BACSU	Acc# P20691
Description				L,,-	
(5-ENOLPYRUVYLSHI)	KIMATE-3	- РНОЅРНА	TE SYNTHASE	(EPSP SYNTHASE)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
25025987_c2_232	1671	3591	207	624 484	4.5e-46
					L
Protein name				Locus Name sp:YRBH_ECOLI	Acc# P45395
Protein name  Description		<u> </u>	J ()		
	KD PROT	EIN IN M	URA-RPON IN	sp:YRBH_ECOLI	P45395
Description	KD PROT	EIN IN M	URA-RPON IN  NT  Length	sp:YRBH_ECOLI	P45395
Description HYPOTHETICAL 35.2			NT	Sp:YRBH_ECOLI  STERGENIC REGION  AA Score	P45395
Description HYPOTHETICAL 35.2 ORF Name	NTID	AAID	<u>NT</u> Length	Sp:YRBH_ECOLI  TERGENIC REGION  AA  Length  Score	P45395  (O328)  Probability
Description  HYPOTHETICAL 35.2  ORF Name  25431625_c3_251	NTID	AAID	<u>NT</u> Length	Sp:YRBH_ECOLI  TERGENIC REGION  AA Length Score  321 224 Locus Name	P45395  (O328)  Probability  1.6e-18  Acc#
Description  HYPOTHETICAL 35.2  ORF Name  25431625_c3_251  Protein name	NTID 1672	<u>AAID</u> 3592	NT Length	Sp:YRBH_ECOLI  TERGENIC REGION  AA Score Length  224  Locus Name  Sp:IHFB_ERWCH	P45395  (O328)  Probability  1.6e-18  Acc#
Description  HYPOTHETICAL 35.2  ORF Name  25431625_c3_251  Protein name  Description	NTID 1672	<u>AAID</u> 3592	NT Length	Sp:YRBH_ECOLI  TERGENIC REGION  AA Score Length  224  Locus Name  Sp:IHFB_ERWCH	P45395  (O328)  Probability  1.6e-18  Acc#
Description  HYPOTHETICAL 35.2  ORF Name  25431625_c3_251  Protein name  Description  INTEGRATION HOST	NTID 1672 FACTOR B	AAID 3592 ETA-SUBU	NT Length 106 NIT (IHF-BE	Sp:YRBH_ECOLI  TERGENIC REGION  AA Score  Length  224  Locus Name  Sp:IHFB_ERWCH  AA Score	P45395  (0328)  Probability  1.6e-18  Acc# P37983
Description  HYPOTHETICAL 35.2  ORF Name  25431625_c3_251  Protein name  Description  INTEGRATION HOST  ORF Name	NTID  1672  FACTOR B  NTID  1673	AAID  [3592  ETA-SUBU  AAID  [3593	NT Length 106 NIT (IHF-BE NT Length	Sp:YRBH_ECOLI  TERGENIC REGION  AA Score  Length 224  Locus Name  Sp:IHFB_ERWCH  ETA)  AA Score	P45395  (O328)  Probability  1.6e-18  Acc# P37983  Probability

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
25564402_c3_285	1674 3594	739	2220 82	9.2e-06
Protein name hypothetical prot	ein SCI7.24c		Locus Name	Acc# T36920
Description				
ORF Name 26359451 c2 249	NTID AAID	NT Length 713	<u>AA</u> <u>Length</u> <u>Score</u>   2142   2272	Probability
Protein name			Locus Name  Sp:UVRB PSEAE	Acc#
Description				P72174:P72 147
EXCINUCLEASE ABC	SUBUNIT B			
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
26750090_c2_245	1676 3596	347	1044 880	4.9e-88
Protein name			Locus Name	Acc#
			sp:PYRD_SALTY	P25468
Description			sp:PYRD_SALTY	P25468
Description (DHODEHASE)			sp:PYRD_SALTY	P25468
	NTID AAID	NT Length	Sp:PYRD_SALTY  AA Length Score	P25468  Probability
(DHODEHASE)	NTID AAID		AA Score	
(DHODEHASE)  ORF Name		Length	AA Score	Probability 4.4e-32 Acc#
(DHODEHASE)  ORF Name  2923562_c2_233	1677 3597	Length	AA Score  Length 352  Locus Name	Probability 4.4e-32 Acc#
ORF Name  2923562_c2_233  Protein name  Description	1677 3597	<u>Length</u>	AA Score Length 352 Locus Name Sp:YRBI_ECOLI	Probability 4.4e-32 Acc# P45396:P45
ORF Name  2923562_c2_233  Protein name  Description		<u>Length</u>	AA Score Length 352 Locus Name Sp:YRBI_ECOLI	Probability 4.4e-32 Acc# P45396:P45
(DHODEHASE)  ORF Name  2923562_c2_233  Protein name  Description  HYPOTHETICAL 20.0	1677 3597  KD PROTEIN IN MUR	Length 177  A-RPON IN NT	AA Score Length 352 Locus Name Sp:YRBI_ECOLI TERGENIC REGION AA Score	Probability  4.4e-32  Acc#  P45396:P45 398
(DHODEHASE)  ORF Name  2923562_c2_233  Protein name  Description  HYPOTHETICAL 20.0  ORF Name	1677   3597	Length  177  A-RPON IN  NT  Length	AA Length 534 Score Locus Name Sp:YRBI_ECOLI TERGENIC REGION AA Length Score	Probability  Acc# P45396:P45 398  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30173201_f2_94	1679	3599	66	201		
Protein name				Locus	Name	Acc#
Description						
NO-HIT	<del></del>					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
30469092_c1_151	1680	3600	250	753	152	6.4e-09
Protein name	-			Locus		Acc#
unknown				gp:MLC1	1622	Z95398
Description						
Mycobacterium lepra	e cosm	id L622.				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	<u>score</u>	Probability
30600453_f3_139	1681	3601	697	2094	777	2.0e-86
Protein name				Locus	Name	Acc#
hypothetical protei	n b2324			pir:B6	5005	B65005
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	Probability
30720027_f3_141	1682	3602	154	465	382	2.9e-35
Protein name				Locus	Name	Acc#
hypothetical protei	n			gp:PPPA	L1	X74218
Description				<u> </u>		
Pseudomonas putida	ruvB, t	colQ, tolk	, tolA, t	olB and op	rL gene	es.
ORF Name			NT	<u>AA</u>	core	Probability
	NTID	<u>AAID</u>	<u>Length</u>	Length -	core	FIODADITICY
31800280_c1_158	NTID 1683	<u>AAID</u>			649	1.5e-63
31800280_c1_158 Protein name			Length	Length =	649	
	1683		Length	Length 918	649 Name	1.5e-63
Protein name	1683		Length	Length 918 Locus	649 Name	1.5e-63 <u>Acc#</u>

ORF Name	NTID	AAID	<u>NT</u> Length	AA Sco	re Probability
31828211_f2_69	1684	3604	1208	3627 29	34 0.0
Protein name				Locus Na	me Acc#
proline dehydroge	enase			gp:ATU392	63 U39263
Description					
Agrobacterium tur (prp) genes, comp		plasmid	pAtR10 pro	line dehydro	genase(putA) and Prp
ORF Name	NTID	AAID	<u>NT</u> Length	AA Scot	re <u>Probability</u>
33229667_c3_270	1685	3605	72	219	
Protein name				Locus Na	me Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Scor	re Probability
33985930_f1_23	1686	3606	288	867	
Protein name				Locus Na	me Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Sco	re Probability
34062503_c1_178	1687	3607	181	546 22	4 1.6e-18
Protein name				Locus Na	<del></del>
Description					
HYPOTHETICAL PRO	rein Hill	49 PRECU	RSOR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Scor	re Probability
34172883_c1_176	1688	3608	166	501 29	6 3.8e-26
Protein name				Locus Na	
Description					
HYPOTHETICAL PRO	rein Hioo	55 PRECU	RSOR		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34409658_f2_84	1689	3609	360	1083	757	5.3e-75
Protein name  carboxyl esterase				Locus pir:S	s <u>Name</u> 57530	<u>Acc#</u> S57530
Description						<del></del>
ORF Name 35157165 cl 191	NTID	<u>AAID</u>	NT Length	AA Length 615	Score	Probability 2.7e-29
	] [1030	] [5010	204			
Protein name methylated-DNApro	-	ysteine		Locus pir:D	8 Name 64604	Acc# D64604
Description			-	_		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36072135_c3_295	1691	3611	686	2061	564	1.5e-54
Protein name					s Name	Acc#
Description				sp:EX	5A_ECOLI	P04993:Q59 378
ALPHA CHAIN)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36112900_f2_91	1692	3612	103	312	253	1.4e-21
<u>Protein name</u>					3 Name U24202	Acc# U24202
Description						
Escherichia coli E (yciA), membrane procardiolipin synthas	otein (	tonB), (yo	iI), puta	tive pota	_	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36129676_c3_252	1693	3613	139	420	85	0.00086
Protein name				Locus	s Name	Acc#
hypothetical prote	in vrvD			pir:G	caaan	G69980
· · · · · · · · · · · · · · · · · · ·		· · ·		] P11.6		909980

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	
3915943_c2_226	1694	3614	414	1245	1163	5.0e-118	
Protein name					s Name TZ_PSEAE	Acc# P55218	
Description							
O-SUCCINYLHOMOSER	RINE SULF	HYDRYLASE,	(OSH SUL	FHYDRYLA	SE)		٦
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	Score	Probability	
3922193_£2_50	1695	3615	381	1146	708	8.3e-70	
Protein name				Locus	s Name	Acc#	
probable pvdS pro	tein			pir:B	70591	B70591	
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability	
3933437_c2_229	1696	3616	202	609	387	8.6e-36	
Protein name				Locus	s Name	Acc#	
hypothetical prot	ein jhp08	367		pir:B	71879	B71879	
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability	
4016943_f2_67	1697	3617	473	1422	653	5.6e-64	
Protein name					s Name	Acc#	
				sp:Y4	WA_RHISN	P55679	
Description							
HYPOTHETICAL ZINC	PROTEAS	E Y4WA,					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability	_
4103293_c1_179	1698	3618	245	738	822	6.9e-82	
Protein name  putative ABC tran	sporter 7	ATP-bindin	g protein		Name 013987	Acc# AF01398	 7
<u>Description</u>				J <u></u>			
Vibrio cholerae s sigma54 (rpoN), pu IIA protein (ptsN)	tative s	igma54 mod	lulation p	_			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4114702_c1_159	1699	3619	119	360	194	2.4e-15
Protein name probable dihydroned	pterin	aldolase,		Locus pir:H	Name 55093	Acc# H65093
Description						·
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4489463_f2_90	1700	3620	425	1278		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4689693_c3_278	1701	3621	372	1119	509	8.2e-55
Protein name				-	Name AA HAEIN	Acc# P44495
Description						
(IPP TRANSFERASE)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4772050_£2_86	1702	3622	441	1326	487	2.2e-46
Protein name					Name BE_HAEIN	<u>Acc#</u> P43745
Description						
DNA POLYMERASE III,	EPSILO	ON CHAIN,				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4816513_c3_294	1703	3623	1318	3957	230	3.1e-41
Protein name					Name B ECOLI	Acc# P08394
Description						
BETA CHAIN)						

ORF Name NTID AA	ID <u>NT</u> Length	AA Score	Probability
4863458_c2_234 1704 3	624 172	519	
Protein name		Locus Name	Acc#
Description			
ио-ніт	*		
ORF Name NTID AA	ID <u>NT</u> Length	AA Length Score	<u>Probability</u>
4876525_c3_293 [1705] 3	625 229	690	
Protein name		Locus Name	Acc#
Description			
ро-ніт			
ORF Name NTID AA	ID NT Length	AA Length Score	Probability
4878135_c2_250 [1706]	626 350	1053 810	1.3e-80
Protein name		Locus Name	Acc#
yhdG homolog		gp:AF040378	AF040378
Description			
Serratia marcescens ribosomal partial cds; and yhdG homolog complete cds.	_	_	
ORF Name NTID AA	ID <u>NT</u> Length	AA Score	Probability
4881700_c3_290 1707 3	479	1440 381	3.7e-35
Protein name hypothetical protein 5		Locus Name	Acc# T00101
Description			
ORF Name NTID AA.	ID <u>NT</u> Length	AA Score	Probability
	233	<u> </u>	
Protein name hypothetical protein		Locus Name gp:AF031940	<del></del>
Description		7 2	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5086693_c3_277	1709	3629	419	1260	793	8.2e-79
Protein name hypothetical protei	n slr00	049		Locus pir:57	Name 74347	Acc# S74347
Description						
ORF Name	NTID	AAID	NT Length	Length	Score	Probability
5098937_f2_51	1710	3630	543	1632	520	3.1e-53
Protein name probable exodeoxyri subunit Description	bonucle	ease VII	Large	Locus pir:C7	Name 75549	<u>Acc#</u> C75549
ORF Name 5110963_c1_162 Protein name	NTID 1711	<u>AAID</u>   3631	<u>NT</u> <u>Length</u>	Length 1680	Score 1056	Probability  1.1e-106  Acc#
Description				sp:ODF	P2_PSEAE	Q59638
COMPLEX, (E2)						<u> </u>
ORF Name	NTID	AAID	NT Length	AA Length [834	Score	Probability [2.4e-33]
Protein name	1712	3632	277	Locus	Name M_HAEIN	Acc# P71396
Description				-		
PUTATIVE FERREDOXII	N-LIKE	PROTEIN H	11684			
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	Length	Score	Probability
5323750_£3_108	1713	3633	104	315		
Protein name				Locus	Name	Acc#
Description						
NO-HIT						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
6484691_f1_26	1714	3634	377	1134 738	5.5e-73
Protein name				Locus Name sp:CYSP_ECOLI	Acc# P16700
Description				<del></del>	
THIOSULFATE-BIND	ING PROTE	IN PRECUE	RSOR		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
806512_c3_279	1715	3635	137	414 171	1.8e-12
Protein name				Locus Name	Acc#
polysialic acid	capsule ex	cpression	protein	pir:B70434	B70434
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
11562_c3_7	1716	3636	78	237	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
20395432_c2_6	1717	3637	70	213	
Protein name				Locus Name	Acc#
Description					·
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
36117135_£1_1	1718	3638	335	1008 1334	3.8e-136
Protein name				Locus Name	Acc#
malate dehydrogen	nase			gp:AF109682	AF109682
Description					<del></del>
Aquaspirillum ar	cticum ma.	late dehy	drogenase	(MDH) gene, comp.	Letecds.

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6682962 <u>f</u> 1_2	1719	3639	85	258		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT			· · · · · · · · · · · · · · · · · · ·		· · · · · · · · · · · · · · · · · · ·	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
13958403_f1_1	1720	3640	399	1200	1251	2.4e-127
Protein name					s Name	Acc# P75802
Description						
HYPOTHETICAL 49.6	KD PROT	EIN IN MOE	EA-DACC IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
16506250_f2_4	1721	3641	140	423	236	4.8e-19
Protein name	_			Locu	s Name	Acc#
unknown	_	<del></del> -		gp:AF	026544	AF026544
Description						
Ralstonia eutropha unknown genes.	phbF a	nd beta-ke	etothiolas	se (bktB)	genes,co	omplete cds; and
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20782550_t3_16	1722	3642	242	729	933	1.2e-93
Protein name					s Name	Acc#
				sp:MT	NG_NEIGO	P08455
Description						
METHYLTRANSFERASE	NGOPII)	(M.NGOPII	.)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24328950_£2_8	1723	3643	153	462	336	2.2e-30
Protein name	•				s Name FH_ECOLI	Acc# P45802
Description				55.11		
HYPOTHETICAL 15.5	KD PROTI	EIN IN MRC	A-PCKA IN	TERGENIC	REGION (	0133)

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
29859790_c2_32	1724	3644	74	225		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
3942592_f2_6	1725	3645	252	759	741	2.6e-73
Protein name hypothetical protein	ein, 26K			Locus	S Name	Acc# JC5479
Description	·			ــــــا لـ		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4103390_f3_15	1726	3646	85	258	323	5.2e-29
Protein name					S Name NG_NEIGO	Acc# P08455
Description				<u> </u>		
METHYLTRANSFERASE	NGOPII)	(M.NGOPII	)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
42837_f1_2	1727	3647	71	216	170	2.5e-12
Protein name					Name	Acc# P08455
Description				<u> </u>		
METHYLTRANSFERASE	NGOPII)	(M.NGOPII	)			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4976512_t2_7	1728	3648	518	1557	1265	7.8e-129
Protein name				Locus	Name	Acc#
threonine dehydra	ase, bio	synthetic		pir:E	75502	E75502
Description						

ORF Name	NTID	<u>AAID</u>	NT Length	<u>AA</u> Length	Score	Probability
7038307_f1_3	1729	3649	127	384	228	6.1e-19
Protein name				-	Name	Acc#
Description						P24666:Q16 035:Q16725
(EC 3.1.3.48) (AD)	POCYTE	ACID PHOSE	PHATASE, I	SOZYME AI	JPHA)	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30175950_f1_1	1730	3650	77	234	292	5.8e-25
Protein name					Name	Acc#
Description				<u> </u>		P45740:P71
THIAMINE BIOSYNTHE	SIS PRO	TEIN THIC				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
4470181_t3_5	1731	3651	156	471	635	4.5e-62
Protein name					Name C_ECOLI	Acc# P30136
Description						
THIAMINE BIOSYNTHE	SIS PRO	TEIN THIC		-		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
7119001_f2_4	1732	3652	73	222	85	0.013
Protein name				· ·	Name 1 HAEIN	Acc#
<u>Description</u>				sp:ra:	T_HABIN	Q57180:005 043
HYPOTHETICAL ABC T	RANSPOR	TER ATP-BI	NDING PRO	TEIN HIIC	51	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24254702_f3_3	1733	3653	252	759	253	1.4e-21
Protein name					Name T_ECOLI	Acc# P37681
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
25787500_c3_6	1734	3654	278	837 503	4.4e-48
Protein name				Locus Name	Acc# GO P72080
Description					
BACTERIOFERRITIN	Α				
ORF Name	NTID	AAID	NT Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
5133575_c3_7	1735	3655	162	489 489	1.3e-46
Protein name				Locus Name sp:BFRB_NEI	Acc# P77914
Description					
BACTERIOFERRITIN	B (BFR A)	(BFR B)			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
21984375_c1_10	1736	3656	473	1422 708	8.3e-70
Protein name				Locus Name	Acc# FT P46681
Description					
ACTIN INTERACTING	PROTEIN	2	·····		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23535910_f1_2	1737	3657	492	1479 1489	1.4e-152
Protein name				Locus Name	Acc#
Description				sp:YEGQ_ECO	P76403:008 007:008010
PUTATIVE PROTEASE	YEGQ,				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
25677176_f3_6	1738	3658	205	615 516	1.8e-49
Protein name site-specific DNA (cytosine-specific				Locus Name	Acc# A64660
Description					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3926262_c2_13	1739	3659	225	678	134	8.6e-08
Protein name				_	s Name 168355	<u>Acc#</u> AF168355
<u>Description</u>			·	_ <u></u>	***	
Proteus mirabilis gene.	telluri	te resist	ance locus	, comple	te sequei	nce; and unknown
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3946943_f1_1	1740	3660	510	1533	808	2.1e-80
Protein name OprM					s Name 011381	Acc# AB011381
Description						
Pseudomonas aerugi	nosa ge	ne for Op	orM, comple	ete cds.		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2110657_c2_3	1741	3661	221	663	570	3.5e-55
Protein name					s Name 26_SYNY3	Acc# P72872
Protein name  Description						
	KD PROT	EIN SLLOS	926			
Description	KD PROT	EIN SLLOS AAID	NT Length			
Description HYPOTHETICAL 37.9			NT	sp: Y9	26_SYNY3	P72872
Description  HYPOTHETICAL 37.9  ORF Name  16040887_f3_11  Protein name	NTID	AAID	<u>NT</u> Length	Sp:Y9  AA  Length  1512  Locus	Score  2557 S Name	P72872 Probability  9.6e-266 Acc#
Description  HYPOTHETICAL 37.9  ORF Name  16040887_f3_11  Protein name  unknown	NTID	AAID	<u>NT</u> Length	Sp:Y9  AA  Length  1512  Locus	26_SYNY3 Score	Probability  9.6e-266
Description  HYPOTHETICAL 37.9  ORF Name  16040887_f3_11  Protein name unknown  Description	NTID 1742	<u>AAID</u> 3662	NT Length	sp:Y9  AA Length 1512 Locus gp:AF	Score 2557 s Name 039312	P72872  Probability  9.6e-266  Acc# AF039312
Description  HYPOTHETICAL 37.9  ORF Name  16040887_f3_11  Protein name  unknown	NTID 1742	AAID  3662  ain 4223	NT Length 504 transferri	AA Length 1512 Locus gp:AF	Score  2557 S Name 039312	P72872  Probability  9.6e-266  Acc# AF039312
Description  HYPOTHETICAL 37.9  ORF Name  16040887_f3_11  Protein name  unknown  Description  Moraxella catarrha	NTID 1742	AAID  3662  ain 4223	NT Length 504 transferri	AA Length 1512 Locus gp:AF	Score  2557 S Name 039312	P72872  Probability  9.6e-266  Acc# AF039312
Description  HYPOTHETICAL 37.9  ORF Name  16040887_f3_11  Protein name  unknown  Description  Moraxella catarrhatransferrin binding	NTID  1742  Alis stra	AAID  3662  ain 4223  n B (tbpE	NT Length 504 transferri s) genes, c	AA Length 1512 Locus gp:AF  n binding	Score  2557  S Name 039312  g proteinds; and u	Probability  9.6e-266  Acc# AF039312  A(tbpA) and anknown gene.
Description  HYPOTHETICAL 37.9  ORF Name  16040887_f3_11  Protein name  unknown  Description  Moraxella catarrhatransferrin binding	NTID  1742  Alis stra g protein  NTID  1743	AAID  3662  ain 4223 n B (tbpE  AAID  3663	NT Length  504  transferri s) genes, c NT Length  108	AA Length 1512 Locus gp:AF  n binding completect AA Length 327	Score  2557  S Name 039312  g proteinds; and u Score  92  Name	Probability  9.6e-266  Acc# AF039312  A(tbpA) and anknown gene.  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	bility
4484567_f1_1	1744	3664	899	2700	4565	0.0	
Protein name transferrin binding	protei	n A		_	s Name 039312		Acc# AF039312
Description					<del>-</del>		
Moraxella catarrhal transferrin binding						_	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
4775207_f2_9	1745	3665	173	522	728	1.6e-	71
Protein name				Locu	s Name		Acc#
transferrin binding	protei	n A		gp:AF	039315		AF039315
Description							
Moraxella catarrhal transferrin binding							
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
33380275_f1_2	1746	3666	65	198			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT			··				
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probab	
35361043_c1_7	1747	3667	62	189	93	0.0004	18
Protein name				Locus	s Name		Acc#
phosphate-binding p phosphate-repressibl				pir:I	64120		I64120
Description					•		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
36501561_c3_9	1748	3668	301	903	842	5.2e-8	34
Protein name					s Name		Acc#
Description				sp:PS	TC_HAEIN		P45191
PHOSPHATE TRANSPORT	SYSTEM	M PERMEASE	PROTEIN	PSTC		_	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5960433_c1_8	1749	3669	60	183		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT			<del></del>	<u> </u>		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4429510_t1_1	1750	3670	477	1434	1328	1.7e-135
Protein name					s <u>Name</u> NB_SALMO	Acc# Q01411
Description				<u></u>		
PHOSPHOMANNOMUTASE,	(PMM)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4459376_c2_15	1751	3671	294	885	575	1.0e-55
Protein name				Locus	s Name	Acc#
Protein name  conserved hypotheti	cal pro	otein	<del> </del>	Locus pir:D		<u>Acc#</u> D75311
	cal pro	otein	:			<del></del>
conserved hypotheti	cal pro	AAID	NT Length			<del></del>
conserved hypotheti				pir:D	75311	D75311
Conserved hypotheti Description  ORF Name  10429517_c1_34  Protein name	NTID 1752	<u>AAID</u>	Length	AA Length	75311 Score	D75311 Probability
Conserved hypotheti Description  ORF Name  10429517_c1_34	NTID 1752	<u>AAID</u>	Length	AA Length	Score  [573] S Name	D75311  Probability  1.7e-55
Conserved hypotheti Description  ORF Name  10429517_c1_34  Protein name	NTID 1752	<u>AAID</u>	Length	AA Length 1242	Score  [573] S Name	D75311  Probability  1.7e-55  Acc#
Conserved hypotheti  Description  ORF Name  10429517_c1_34  Protein name  conserved hypotheti	NTID 1752	<u>AAID</u>	Length	AA Length 1242 Locus pir:A	Score  [573] S Name	D75311  Probability  1.7e-55  Acc#
Conserved hypotheti  Description  ORF Name  10429517_c1_34  Protein name  Conserved hypotheti  Description	NTID 1752 cal pro	AAID 3672 stein	Length 413	AA Length 1242 Locus pir:A	Score [573] S Name 75525	D75311  Probability  1.7e-55  Acc# A75525
Conserved hypotheti  Description  ORF Name  10429517_c1_34  Protein name  Conserved hypotheti  Description  ORF Name	NTID  1752  cal pro	AAID 3672 Stein AAID	Length 413  NT Length	AA Length Locus pir:A  AA Length 1242  Locus pir:A	Score [573] S Name 75525	Probability  1.7e-55  Acc# A75525  Probability
Conserved hypotheti  Description  ORF Name  10429517_c1_34  Protein name  Conserved hypotheti  Description  ORF Name  12384625_c3_48	NTID  1752  cal pro	AAID 3672 Stein AAID	Length 413  NT Length	AA Length Locus pir:A  AA Length 1242  Locus pir:A	Score 573 S Name 75525 Score 386 S Name	Probability  1.7e-55  Acc# A75525  Probability  1.1e-35  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
15915625_f2_13	1754	3674	170	513	
Protein name				Locus Name	Acc#
Description					
по-ніт					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
21673425_f2_12	1755	3675	447	1344 507	1.7e-48
Protein name				Locus Name	Acc#
Description				sp:UBIH_ECOLI	P25534
UBIH PROTEIN,					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
2195931_c1_29	1756	3676	69	210 93	0.0018
Protein name				Locus Name	Acc#
conserved hypothe	tical pro	otein aq_:	2107	pir:F70480	F70480
Description					-
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
22355001_f3_23	1757	3677	73	222	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Score	Probability
23954035_c3_50	1758	3678	175	528 119	3.8e-05
Protein name				Locus Name	Acc#
conserved hypothe	tical pro	tein aq_	2107	pir:F70480	F70480
Description					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
25665885_c2_36	1759	3679	241	726 389	6.3e-45
Protein name				Locus Name sp:MIAE_SALTY	<u>Acc#</u> Q08015
Description			,	•	
TRNA- (MS [2					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
30100880_c3_51	1760	3680	201	606 186	1.7e-14
Protein name				Locus Name	Acc#
hypothetical protei	in aq_2:	L08		pir:G70480	G70480
Description					
ORF Name	NTID	AAID	NT Length	AA Length Score	Probability
3994052_f1_3	1761	3681	192	579 682	4.7e-67
•					
Protein name				Locus Name	Acc#
Protein name probable dctp deam	nase			Locus Name	<u>Acc#</u> B71565
	inase			_	<del></del>
probable dctp deams  Description  ORF Name	NTID	AAID	NT Length	pir:B71565  AA Length Score	B71565  Probability
probable dctp deami		<u>AAID</u> 3682		pir:B71565	B71565
probable dctp deams  Description  ORF Name  4109790_c1_28  Protein name	NTID 1762	3682	Length 155	pir:B71565  AA Length Score	B71565  Probability
probable dctp deams  Description  ORF Name  4109790_c1_28	NTID 1762	3682	Length 155	pir:B71565   AA   Score   468   176	B71565  Probability  2.2e-12
probable dctp deams  Description  ORF Name  4109790_c1_28  Protein name	NTID 1762	3682	Length 155	pir:B71565  AA Score Length 468 176  Locus Name	Probability  2.2e-12  Acc#
probable dctp deams  Description  ORF Name  4109790_c1_28  Protein name  conserved hypothetic	NTID 1762	3682	Length 155 2107 NT	pir:B71565  AA Score Length 468 176  Locus Name pir:F70480  AA Score	Probability  2.2e-12  Acc#
probable dctp deams  Description  ORF Name  4109790_c1_28  Protein name  conserved hypothetic  Description	NTID 1762 Cal pro	3682 Stein aq	Length [155] 2107	pir:B71565  AA Score Length 176  Locus Name pir:F70480	Probability  2.2e-12  Acc# F70480
probable dctp deams  Description  ORF Name  4109790_c1_28  Protein name  conserved hypothets  Description  ORF Name	NTID  1762  cal pro	3682 Stein aq	Length  155  2107  NT Length	AA Score Length 176 Locus Name pir:F70480  AA Score Locus Name Locus Name Length 1482 Locus Name Locus Name	Probability  2.2e-12  Acc# F70480  Probability  3.7e-44  Acc#
probable dctp deams  Description  ORF Name  4109790_c1_28  Protein name  conserved hypothets  Description  ORF Name  4319837_c2_37	NTID  1762  cal pro	3682 Stein aq	Length  155  2107  NT Length	AA Score Length 176  Locus Name pir:F70480  AA Score Length 468  Locus Name pir:F70480	Probability  2.2e-12  Acc# F70480  Probability  3.7e-44

ORF Name	NTID AAID NT Length	AA Length Score	Probability
4345068_f3_21	1764 3684 128	387 177	1.5e-13
Protein name		Locus Name sp:YOHJ_ECOLI	Acc# P33372
Description			
HYPOTHETICAL 14.6	KD PROTEIN IN PBPG-CDD IN	rergenic region	
ORF Name	NTID AAID NT Length	AA Length Score	Probability
4790637_±3_22	1765 3685 183	552 295	4.8e-26
Protein name		Locus Name sp:YOHK_HAEIN	Acc# P45146
Description		<del></del>	
HYPOTHETICAL PRO	TEIN HI1298		
ORF Name	NTID AAID NT Length	AA Length Score	Probability
5900203_f1_1	1766 3686 689	2070 1609	2.8e-165
Protein name		Locus Name sp:REP_ECOLI	Acc# P09980
Description			
ATP-DEPENDENT DNA	A HELICASE REP,		· · · · · ·
ORF Name	NTID AAID <u>NT</u> Length	AA Length Score	Probability
6658527_c2_39	1767 3687 184	555 228	5.6e-18
Protein name		Locus Name	Acc#
L	etical protein aq_2107	pir:F70480	F70480
<u>Description</u>			
ORF Name	NTID AAID NT Length	Length	Probability
7226518_£2_17	1768 3688 102	309 112	1.2e-06
Protein name hypothetical prot	ein	Locus Name gp:POL010393	Acc# AJ010393
Description	· · · · · · · · · · · · · · · · · · ·	<u> </u>	
Pseudomonas oleo	vorans phal and phar genes,	and ORF1, ORF2 (par	tial) and

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
22897332_c2_15	1769	3689	153	462	338	1.3e-	30
Protein name					s Name AH_BACNO		Acc# P04953
Description							
SUBUNITS PILIN)							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
36210875_f2_3	1770	3690	883	2652	3272	0.0	
Protein name					Name		Acc#
Description							P36683:P36 648:Q59382 :P75652
(ACONITASE 2)			<u> </u>				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
14853143_c1_9	1771	3691	703	2112	1460	1.7e-	149
Protein name		-			S Name GF_NEIME		<u>Acc#</u> Q51152
Description							
HYPOTHETICAL 83.1 k	D PROTE	IN IN REG	ION E			<del> </del>	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
16050817_f1_1	1772	3692	225	678	203	2.7e-	16
Protein name				Locus	s Name		Acc#
hypothetical protei	n s1107	88		pir:S	77018		S77018
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
10175877_£3_73	1773	3693	264	795	124	2.2e-	11
Protein name DnrD protein					Name 1131715		Acc# AJ131715
Description			<del></del>	J [			
Pseudomonas stutzer	i dnrD	gene and (	ORF194 (p	artial) a	and ORF63	(parti	al).

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
10195250_f2_49	1774	3694	81	246	
Protein name		_	-	Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
10546930_f1_18	1775	3695	239	720 576	8.1e-56
Protein name				Locus Name	Acc# N P45322
Description					
MOLYBDENUM TRANSPO	RT SYST	EM PERMEAS	SE PROTEIN	MODB	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
11113152_f3_70	1776	3696	142	429 205	1.7e-16
Protein name				Locus Name	Acc#
hypothetical prote	in APET	291		pir:D72603	D72603
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
12367711_f3_63	1777	3697	1 259		G
		1 1	233	780 438	3.4e-41
Protein name			259	Locus Name sp:MODD_AZOV	Acc#
Protein name  Description	<b>-</b>	J L	239	Locus Name	Acc#
	RT ATP-	J L		Locus Name sp:MODD_AZOV	Acc#
Description	ORT ATP-	J L		Locus Name sp:MODD_AZOV	Acc#
Description  MOLYBDENUM TRANSPO		BINDING PR	ROTEIN MOD	Locus Name  sp:MODD_AZOV	Acc# P37732
Description  MOLYBDENUM TRANSPO	NTID	BINDING PR	ROTEIN MODE  NT  Length	Locus Name  sp:MODD_AZOV  AA Length Score	Acc# P37732  Probability
Description  MOLYBDENUM TRANSPO  ORF Name  15710327_c1_84	<u>NTID</u>	BINDING PR	ROTEIN MODE  NT  Length	Locus Name  sp:MODD_AZOV  AA Length  801  427	Acc# P37732  Probability 5.0e-40
Description  MOLYBDENUM TRANSPO  ORF Name  15710327_c1_84  Protein name	<u>NTID</u>	BINDING PR	ROTEIN MODE  NT  Length	Locus Name  sp:MODD_AZOV  D  AA Length 801 427  Locus Name	Acc# P37732  Probability  5.0e-40  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
15781576_c2_103	1779	3699	223	672	594	1.0e-	57
Protein name					S Name		Acc#
Description							P36857:P75 656
HYPOTHETICAL 25.1	KD PROTE	IN IN HPT	-PAND INT	ERGENIC I	REGION		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
19735188_£3_58	1780	3700	677	2034	484	3.8e-	70
<u>Protein name</u>					s Name		Acc#
nitrate/nitrite se	nsory pr	otein		gp:PS	NARXL		Y15252
Description							
Pseudomonas aerugi nifM, moaA genes.	nosa nar	X, narL,	narKl, na	rK2, nar	3, narH,	narJ,r	narI,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	<u>bilit</u> y
19806552_f2_31	1781	3701	187	564	134	5.5e-	08
Protein name Notch homolog				_	Name 033013		<u>Acc#</u> AF033013
				_	Name 033013		<u>Acc#</u> AF033013
Notch homolog	homolog	mRNA, par	tial cds.	gp:AF			
Notch homolog  Description	homolog	mRNA, par	tial cds. <u>NT</u> Length	gp:AF		Proba	
Notch homolog  Description  Bombyx mori Notch			NT	gp:AF	033013	Proba	AF033013
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51  Protein name	NTID	AAID	NT Length	gp:AF	Score [142]		AF033013
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51	NTID	AAID	NT Length	gp:AF	033013 <u>Score</u> 142		AF033013
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51  Protein name	NTID	AAID	NT Length	gp:AF	Score [142]		AF033013  bility  09  Acc#
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51  Protein name  Notch homolog	NTID 1782	<u>AAID</u> 3702	NT Length 180	AA Length  543  Locus  gp:AF6	Score [142]		AF033013  bility  09  Acc#
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51  Protein name  Notch homolog  Description	NTID 1782	<u>AAID</u> 3702	NT Length 180	AA Length  543  Locus  gp:AF6	Score [142]	1.3e-	AF033013  bility  09  Acc#
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51  Protein name  Notch homolog  Description  Bombyx mori Notch	NTID 1782 homolog	AAID  3702  mRNA, par	NT Length 180	AA Length  543  Locus  gp:AF	Score 142 S Name 033013	1.3e-	AF033013  bility  09  Acc# AF033013  bility
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51  Protein name  Notch homolog  Description  Bombyx mori Notch  ORF Name	NTID 1782 homolog NTID	AAID  MRNA, par  AAID	NT Length 180 tial cds.	AA Length  Gp:AF  AA Locus  AA Length  AA Length  B10  Locus	Score 142 S Name 033013	1.3e-	AF033013  bility  09  Acc# AF033013  bility
Notch homolog  Description  Bombyx mori Notch  ORF Name  19806552_f3_51  Protein name  Notch homolog  Description  Bombyx mori Notch  ORF Name  20423500_f2_33	NTID 1782 homolog NTID	AAID  MRNA, par  AAID	NT Length 180 tial cds.	AA Length  Gp:AF  AA Locus  AA Length  AA Length  B10  Locus	Score  142 S Name 033013  Score  420 S Name	1.3e-	AF033013  bility  09  Acc#  AF033013  bility  39  Acc#

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20587686_f1_8	1784	3704	139	420		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
20876387_f1_13	1785	3705	234	705	247	5.9e-21
Protein name					s Name	Acc# P32157
Description				<u></u>	<u></u>	
HYPOTHETICAL 26.6	KD PROTI	EIN IN KDO	T-CPXA IN	TERGENIC	REGION	(0234)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21485962_c2_129	1786	3706	63	189		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT	<del>-</del>					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
21673452_c3_141	1787	3707	448	1347	1455	5.8e-149
Protein name				_	s Name	Acc#
nitrate extrusion	protein			gp:PS	NARXL	Y15252
<u>Description</u>						
Pseudomonas aerugi nifM, moaA genes.	lnosa nai	rX, narL,	narK1, na	rK2, nar	G, narH,	narJ,narI,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21688888_c1_76	1788	3708	332	999	1035	1.8e-104
Protein name					s Name	Acc#
Description				ap: In	TI SWIII	P55913:006 955
THIAMINE BIOSYNTHE	ESIS PRO	TEIN THII				

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22000717_£1_19	1789	3709	145	438		
Protein name				Locu	s Name	Acc#
Description						•
NO-HIT					*	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22378418_£1_17	1790	3710	283	852	502	5.6e-48
Protein name				,	s Name DA_HAEIN	Acc# P45323
Description						
MOLYBDATE-BINDING	PERIPLAS	MIC PROTE	IN PRECUR	SOR		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
22554031_£3_60	1791	3711	194	585	499	1.2e-47
Protein name				•	S Name AB_ECOLI	Acc# P30746
Description				•		<del></del>
MOLYBDENUM COFACT	OR BIOSYN	THESIS PR	OTEIN B			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u>	Score	Probability
			<u> </u>	Length		
24068812_f1_12	1792	3712	258	Length 777	561	3.1e-54
Protein name nitrate/nitrite r				777	S Name	3.1e-54 Acc# Y15252
Protein name				777 Locus	S Name	Acc#
Protein name  nitrate/nitrite r	egulatory	protein	258	Locus gp:PS	NARXL	Acc# Y15252
Protein name nitrate/nitrite r  Description Pseudomonas aerug	egulatory	protein	258	Locus gp:PS	NARXL	Acc# Y15252
Protein name  nitrate/nitrite r  Description  Pseudomonas aerug nifM, moaA genes.	egulatory	protein X, narL,	narK1, na	Locus gp:PS rK2, narc	NARXL G, narH,	Acc# Y15252 narJ,narI,
Protein name  nitrate/nitrite r  Description  Pseudomonas aerug nifM, moaA genes.  ORF Name	egulatory Jinosa nar NTID	protein X, narL,	narK1, na  NT Length	Locus gp:PS  rK2, narc  AA  Length	NARXL G, narH,	Acc# Y15252 narJ,narI,
Protein name  nitrate/nitrite r  Description  Pseudomonas aerug nifM, moaA genes.  ORF Name  24423375_f1_5	egulatory Jinosa nar NTID	protein X, narL,	narK1, na  NT Length	Locus gp:PS  rK2, narc  AA  Length	S Name NARXL G, narH, Score	Acc# Y15252 narJ,narI, Probability

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24423375_f2_32	1794	3714	66	201	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24651536_f1_16	1795	3715	200	603 310	1.2e-27
Protein name				Locus Name	Acc#
				sp:Y903_SYNY3	Q55371
Description					
HYPOTHETICAL 16.5	KD PROT	EIN SLR09	03		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
					0.040
25507260_f1_9	1796	3716	338	1017 82	0.048
Protein name		3716	338	Locus Name	. Acc#
		3716	338		
Protein name		3716	338	Locus Name	Acc#
Protein name MHC class I antige		AAID	NT Length	Locus Name	Acc#
Protein name  MHC class I antige  Description	en		NT	Locus Name pir:I57454  AA Score	. <u>Acc#</u> I57454
Protein name  MHC class I antigo  Description  ORF Name  275283_f1_15  Protein name	NTID	AAID 3717	NT Length	Locus Name pir:157454  AA Length Score	Acc# I57454  Probability
Protein name  MHC class I antige  Description  ORF Name  275283_f1_15	NTID	AAID 3717	NT Length	<u>Locus Name</u> pir:I57454  AA Length 591 142	Probability 7.9e-10
Protein name  MHC class I antigo  Description  ORF Name  275283_f1_15  Protein name	NTID	AAID 3717	NT Length	Locus Name pir:I57454  AA Length 591 142 Locus Name	Probability 7.9e-10  Acc#
Protein name  MHC class I antige  Description  ORF Name  275283_f1_15  Protein name  hypothetical protein	NTID	AAID 3717	NT Length	Locus Name pir:I57454  AA Length 591 142 Locus Name	Probability 7.9e-10  Acc#
Protein name  MHC class I antige  Description  ORF Name  275283_f1_15  Protein name  hypothetical protein  Description	NTID 1797 ein Rv24!	AAID 3717	NT Length 196	Locus Name pir:I57454  AA Score Length 591 Locus Name pir:D70864  AA Score	Probability 7.9e-10  Acc# D70864
Protein name  MHC class I antige  Description  ORF Name  275283_f1_15  Protein name  hypothetical protein  Description  ORF Name	NTID 1797 ein Rv249	AAID 3717 33C AAID	NT Length 196	Locus Name pir:I57454  AA Score Length 142 Locus Name pir:D70864  AA Score	Probability 7.9e-10  Acc# D70864
Protein name  MHC class I antige  Description  ORF Name  275283_f1_15  Protein name  hypothetical protein  ORF Name  2853437_f1_11	NTID 1797 ein Rv249	AAID 3717 33C AAID	NT Length 196	Locus Name pir:I57454  AA Score Length 591 142  Locus Name pir:D70864  AA Score	Probability 7.9e-10  Acc# D70864  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
29432768_c2_123	1799	3719	91	276			
Protein name				Locus	Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
30509827_£1_7	1800	3720	134	405			
Protein name				Locus	Name		Acc#
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
31453162_c2_128	1801	3721	77	234	197	1.2e-1	.5
Protein name				Locus	Name		Acc#
hypothetical prote	in			gp:AF	213822		AF213822
Description							
Zymomonas mobilis	strain	ZM4 fosmio	d clone 42	B3, compl	lete sequ	ience.	···
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
33331633_c1_83	1802	3722	521	1566	2362	4.4e-2	45
Protein name							
				Locus	Name		Acc#
respiratory nitrat	e reduc	tase beta	subunit	Locus gp:PSI			<u>Acc#</u> Y15252
Description	e reduc	tase beta	subunit				
				gp:PSI	NARXL	narJ,na	Y15252
Description Pseudomonas aerugi				gp:PSI	NARXL	narJ,na	Y15252
Description  Pseudomonas aerugi nifM, moaA genes.	nosa na	rX, narL,	narK1, na	gp:PSi	NARXL		Y15252
Description  Pseudomonas aerugi nifM, moaA genes.  ORF Name	nosa na	rX, narL,	narK1, na NT Length	gp:PSN rK2, narc  AA Length 525	NARXL		Y15252
Description  Pseudomonas aerugi nifM, moaA genes.  ORF Name  33758515_f1_6	nosa na	rX, narL,	narK1, na NT Length	gp:PSN rK2, narc  AA Length 525	NARXL  , narH,  Score		Y15252 arI, ility

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36351552_£3_61	1804	3724	91	276	145	3.8e-10
Protein name hypothetical prot	ein ssrl	527		_	<u>s Name</u>	Acc# S75710:S75
<u>Description</u>						718
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
36371012_c2_102	1805	3725	60	183		
Protein name				Locu	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
3906555_£2_36	1806	3726	172	519	296	3.8e-26
Protein name				Locu	s Name	Acc#
probable molybden	um-pterin	n-binding	-protein	pir:S	57954	S57954
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4011062_c1_91	1807	3727	427	1284	1152	7.4e-117
Protein name				_	s Name	Acc#
nitrate extrusion	protein			gp:PS	NARXL	Y15252
Description						
Pseudomonas aerug nifM, moaA genes.	inosa na	rX, narL,	narKl, na	rK2, nar	G, narH,	narJ, narI,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4070308_c3_134	1808	3728	442	1329	676	2.0e-66
Protein name					s Name EA_HAEIN	Acc# P45210
Description				<del></del>		
MOLYBDOPTERIN BIO	SYNTHESIS	S MOEA PR	OTEIN			

ORF Name	NTID AAID	<u>NT</u> Length	AA Length	Score	Probability
4344003_t3_59	1809 3729	364	1095	737	7.0e-73
Protein name			Locus	Name A_HAEIN	Acc# P45311
Description					
MOLYBDENUM COFACTOR	BIOSYNTHESIS PR	OTEIN A	<del>- %</del>		
ORF Name 4788876 c3 133	NTID AAID	NT Length 248	AA Length	Core	Probability [2.2e-69]
	1010 3730	240	L		
Protein name respiratory nitrate	reductase gamma	subunit	Locus gp:PSN/		Acc# Y15252
Description					
Pseudomonas aerugin nifM, moaA genes.	osa narX, narL,	narKI, na	rK2, narG,	narH,	narJ,narI,
ORF Name	NTID AAID	<u>NT</u> Length	AA Length	<u>Score</u>	Probability
4797093_f3_52	1811 3731	157	474		
Protein name			Locus	Name	Acc#
Description					
NO-HIT					
ORF Name 4806502_c2_127	NTID AAID	NT Length	AA Length 309	core	Probability 3.5e-07
Protein name  negative regulator  Description	of translation		Locus gp:AF21		<u>Acc#</u> AF213822
Zymomonas mobilis s	train ZM4 fosmid	clone 42	B3, comple	te sequ	ence.

ORF Name	NTID F	AAID	NT Length	<u>AA</u> Length	Score	Probab	ility
4886251_f2_35	1813	3733	168	507	354	2.7e-3	2
Protein name   molybdenum cofactor	hiosymth	esis pro	tein C		Name 108766	<del></del>	<u>Acc#</u> AF108766
Description				] gp:Ar			AF100700
Rhodobacter sphaero	nides Asmi	(AsmA)	gene na	rtial cds	s: YbaU		
(ybaU), anthranilate anthranilatesynthase anthranilatephosphophosphosphotophosphatesynthase (moaC), molybdenum co	synthase componer cibosyltractryC), mol	componen nt II (tr ansferase lybdenum	t I (trp pG), (trpD), cofactor is prote	E), YibQ indole-3 biosynth in A (mos	(yibQ), 3-glycero nesis pro	tein C	
ORF Name	NTID I	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
4897576_c3_147	1814	3734	69	210			
Protein name				Locus	s Name		Acc#
Description							
NO-HIT					-		
ORF Name	NTID 2	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
5282812_f3_56	1815	3735	65	198	52	0.032	
Protein name					<u>Name</u>		Acc#
MDP1				gp:AB	013441		AB013441
Description		- MDD1 - a	loma Loho	ada	<del> </del>		
Mycobacterium bovi	gene roi						
ORF Name	NTID A	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
630053_t2_47	1816	3736	385	1158	320	1.1e-2	8
Protein name ORF396 protein				Locus gp:PS	Name DNGC		<u>Acc#</u> Z73914
Description				<u> </u>			
Pseudomonas stutze	ri orf175	gene.					
ORF Name	NTID 2	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	ility
635903_f1_10	1817	3737	75	228			
Protein name				Locus	s_Name		Acc#
Description							
NO-HIT		<u> </u>					<u> </u>

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
7064692_c1_86	1818	3738	369	1110	415	9.3e-39
Protein name NifM protein				Locu gp:PS	s Name NARXL	Acc# Y15252
Description						
Pseudomonas aerugin nifM, moaA genes.	iosa na	rX, narL,	narK1, na	rK2, nar	G, narH,	narJ,narI,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
7225637_£3_50	1819	3739	1739	5220	540	6.2e-50
Protein name filamentous hemagg		_	ein	Locu pir:T	s <u>Name</u> 09083	Acc# T09083
PspA:probable secret  Description				]		
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	AA Length	Score	Probability
9814751_c3_131	1820	3740	1271	3816	5075	0.0
Protein name alpha-subunit of n	itrate 1	reductase		_	s Name U71398	Acc# U71398
Description				<u> </u>		
Pseudomonas fluore andbeta-subunit (na				pha-subu	nit (nar(	3)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
1176576_f3_28	1821	3741	157	474	314	4.7e-28
Protein name					s Name	Acc#
Description						P52088:P75 703
HYPOTHETICAL 17.0	KD PROT	EIN IN PRO	C-AROL IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
14647507_f2_20	1822	3742	405	1218	386	1.1e-35
Protein name conserved hypothet:	ical pro	otein aq_7	40	Locu pir:A	s Name 70365	<u>Acc#</u> A70365
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
23443752_f3_23	1823	3743	661	1986 298	3.4e-23
Protein name				Locus Name sp:YTRP_PSEPU	Acc# P40604
Description					
HYPOTHETICAL 62.7	KD PROT	EIN IN TRI	PE-TRPG IN	TERGENIC REGION	PRECURSOR
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23610636_c3_58	1824	3744	274	825 755	8.7e-75
Protein name				Locus Name sp:YQCB_HAEIN	Acc# P44197
Description					•
HYPOTHETICAL PROT	EIN HI14	35			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
24644035_c2_45	1825	3745	219	660 255	8.4e-22
Protein name probable citrate	lvase be	ta chain		Locus Name	Acc# T35062
Description				J P11.133002	155002
					•
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
250251_f2_18	1826	3746	192	579 525	2.0e-50
Protein name				Locus Name sp:PUR6_HAEIN	Acc# P43849
Description					
(EC 4.1.1.21) (AI	R CARBOX	YLASE) (A	IRC)		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
25485763_f1_1	1827	3747	89	270	
Protein name				Locus Name	Acc#
Description					
NO-HIT	<u></u>				· · · · · · · · · · · · · · · · · · ·

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
25510974_c1_34	1828	3748	185	558	331	7.4e-30
Protein name					S Name EQ_ECOLI	Acc# P77234
Description						···
HYPOTHETICAL 37.3	KD PROT	EIN IN LE	US-GLTL IN	TERGENIC	REGION	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
29304668_f3_30	1829	3749	302	909	556	1.1e-53
Protein name				Locus	s Name	Acc#
Description				sp:SY	K3_ECOLI	P03812:P78
HYPOTHETICAL LYSYL	-TRNA S	YNTHETASE	HOMOLOG,	(GX)		
ORF Name	NTID	AAID	<u>NT</u> Lengt.h	<u>AA</u> Length	Score	Probability
31275301_f1_11	1830	3750	352	1059	516	1.8e-49
Protein name					Name KB_ECOLI	<u>Acc#</u> P05459
Description					•	
ERYTHRONATE-4-PHOS	PHATE D	EHYDROGEN	ASE,	· · · · · · · · · · · · · · · · · · ·		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
34179211_f3_24	1831	3751	330	993	158	1.2e-08
Protein name probable protein sephosphatase	erine-th	nreonine		Locus pir:C	Name 75297	Acc# C75297
Description						
ORF Name 36365625_f2_17	NTID 1832	AAID 3752	NT Length	AA Length 444	Score	Probability
Protein name				Locus	Name	Acc#
hypothetical prote	in jhp13	377		pir:D7	71815	D71815
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5118952_c1_35	1833	3753	403	1212	703	2.8e-69
Protein name					s Name R2_PSEAE	Acc# Q51551
Description				<del></del>	· · · · · · · · · · · · · · · · · · ·	
CATALYTIC CHAIN)						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
5275300_f3_22	1834	3754	347	1044	954	7.1e-96
Protein name			•	-	s Name OB_ECOLI	Acc# P12996
<u>Description</u>						
BIOTIN SYNTHASE,	(BIOTIN S	YNTHETASE	:)		<del></del>	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
5948342_f3_:27	1835	3755	259	780	530	6.0e-51
<u>Protein name</u>					s Name RK_PSEAE	Acc# P72158
Description						
(AIR CARBOXYLASE)	(AIRC)					
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
7041_f1_8	1836	3756	132	399	155	1.5e-10
<u>Protein name</u>					Name	<u>Acc#</u> 066608
Description						
(AIR CARBOXYLASE)	(AIRC)					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
12991392_f2_18	1837	3757	101	306	160	9.7e-12
Protein name unknown					Name 008856	Acc# U08856
Description				٦ <u>۲.</u>		
Paracoccus denitri	ficans i	nsertion	sequence	IS1248b,	complete	sequence.

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
15110912_c3_69	1838	3758	358	1077	690	6.7e-68
Protein name			_		s Name JM_BACSU	Acc# P54550
Description				<del></del>		
PROBABLE NADH-DEPEN	DENT FI	AVIN OXIL	OREDUCTAS	E YQJM,		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
15632781_c2_61	1839	3759	79	240		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT					-	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
15673816_f3_31	1840	3760	472	1419	1772	1.5e-182
Protein name					s Name	Acc#
type I site-specifi chain R:type I restr	_			pir:J	C5216	JC5216
R:type I restriction	-modifi	cation sy	stem,			
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
19823578_£3_29	1841	3761	405	1218	130	1.0e-07
Protein name				_	s Name	Acc#
hypothetical protei	.n			pir:A	75592	A75592
Description						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21642052_c3_70	1842	3762	232	699	686	1.8e-67
Protein name					s Name	Acc#
<u>Description</u>				sp:YC	78_HAEIN	Q57431:005 050
PUTATIVE NAD(P)H N	TRORED	JCTASE,		<u></u>		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
21673201_±3_25	1843	3763	220	663	364	2.4e-33
Protein name protein Tp70 Description		<del></del>			<u>s Name</u> 71309	Acc# A71309:S18 231:S19826
ORF Name 2189075_c2_66	NTID 1844	<u>AAID</u>	NT Length	AA Length	Score	Probability
Protein name				Locu	s Name	Acc#
Description				sp:YP	RO_OWEFU	P21260:P21 261
HYPOTHETICAL PROI	INE-RICH	PROTEIN	(FRAGMENT)			
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
22143752_f1_9	1845	3765	636	1911	2756	7.9e-287
type I site-speci chain R:type I res R:type I restricti Description	triction	enzyme,	Hsd, chain	pir:J	s Name C5216	Acc# JC5216
ORF Name  23437551_f1_4	NTID 1846	<u>AAID</u>	NT Length	AA Length 252	Score	Probability
Protein name Description				Locus	s Name	Acc#
NO-HIT						
ORF Name 23490937_c3_81	NTID 1847	<u>AAID</u>	NT Length	AA Length 282	Score 73	Probability 0.037
Protein name nicotinamide aden dehydrogenase	ine dinuc	cleotide			S Name 025836	Acc# AF025836
Description						
Echinostoma sp.I. subunit 1 (ND1) ge partial cds.						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabi	lity
24704462_c2_51	1848	3768	206	621	296	3.8e-26	
Protein name  cinnamyl-alcohol de	ehydroge	enase		_	s Name 083333		<u>Acc#</u> F083333
Description							
Medicago sativa cir	nnamyl-a	alcohol de	hydrogena	se (MsaC	adl) mRN	A,complet	e cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabi	lity
32667715_f3_26	1849	3769	188	567	106	0.00093	
Protein name				Locu	s Name	:	Acc#
hypothetical prote	in TP057	0		pir:H	71308	Н	71308
Description							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabi	lity
35350802_f1_5	1850	3770	167	504	345	2.4e-31	
Protein name				Locu	s Name	:	Acc#
putative transposas	se .			gp:AF	007429	A	F007429
Description							
Haemophilus paraga	llinarum	n IS-like	putative	transpos	ase gene,	complete	cds.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabi:	lity
36564842_f2_19	1851	3771	460	1383	573	1.7e-55	
Protein name				Locu	s Name	<u>;</u>	Acc#
type I site-specifichain S:type I restriction  Description	ciction	enzyme, H	sd, chain	pir:J	C5218	J	C5218
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probabil	Lity
4032715_f3_27	1852	3772	116	351	214	1.8e-17	
Protein name					s Name SN_RHISN		<u>Acc#</u> 50358
Description					:=		
HYPOTHETICAL 14.4	KD PROTE	IN Y4SN			<u> </u>		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
4111333_f3_33	1853	3773	201	603	238	5.3e-2	20
Protein name					s Name HR_PSEPU		Acc# P10183
Description							
TRANSCRIPTIONAL A	CTIVATOR	PROTEIN N	AHR				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probal	oility
4895127_c1_36	1854	3774	116	351	334	3.6e-3	30
Protein name					s Name		Acc#
Orf8				gp:AB	011413		AB011413
Description							
Streptomyces grise and complete cds.	eus genes	s for Orf2	, Orf3, C	rf4, Orf	5, AfsA,	Orf8,p	artial
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	oility
7034808_c1_49	1855	3775	70	213	51	0.033	
Protein name hypothetical prote	ein ZK856	5.5		Locus pir:T	s Name 28044		Acc# T28044
Description				J			
ORF Name 7080001_f1_2 Protein name	NTID 1856	<u>AAID</u>	NT Length 371	AA Length 1116 Locus	Score 298 Name	Probab 2.3e-2	
<u>Description</u>				<u> </u>	CG_ECOLI		P55140
HYPOTHETICAL 34.9	KD PROTI	EIN IN CYS	J-ENO INT	ERGENIC	REGION (C	0313)	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	
7083578_c1_37	1857	3777	60	183	162	2.4e-1	.1
Protein name NADP-dependent alo	cohol hyd	irogenase			s Name FL1063		<u>Acc#</u> AL121862
Description				- L			
Leishmania major	Friedlin	chromosom	e 23 cosm	id L1063	, complet	e cds.	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
9782666_£1_7	1858	3778	554	1665	2520	8.0e-262
Protein name ALXA and HSDM				_	S Name U46781	Acc# U46781
Description						<b>:</b>
Pasteurella haemo partial cds, leuko methylase subunit	toxin tra	anscript:	ional activ	ator and	restrict:	ion modification
ORF Name	NTID	<u>AAID</u>	NT Length	AA Length	Score	Probability
Protein name Description		] [3779]			s Name	Acc#
NO-HIT						
ORF Name 16180437_f3_18	NTID 1860	<u>AAID</u>	NT Length	AA Length 1509	<u>Score</u>	Probability 1.7e-149
Protein name  Description					S Name BD_ECOLI	Acc# P25526
(SSDH)				***************************************		
ORF Name  19806552_f1_1	NTID [1861	<u>AAID</u>	NT Length	AA Length 525	Score	Probability 2.6e-07
Protein name				Locus	s Name	Acc#
probable ankyrin			·	pir:H	71274	H71274
Description						
ORF Name	NTID	AAID	NT Length	AA Length	Score	Probability
24407502_f3_17	1862	3782	224	675	392	2.5e-36
Protein name  glycine betaine/catering transporter (membra)  Description			ABC	Locus pir:F	Name 59670	<u>Acc#</u> F69670
<del></del>						

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
25900252_c2_30	1863 3783	3 417	1254	138	1.3e-05
Protein name			Locus	Name	Acc#
putative natural re	sistance-asso	ciated	gp:CCA	133735	AJ133735
Description					
Cyprinus carpio mRI protein (NRAMP).	NA for putativ	ve natural re	esistance-	associat	edmacrophage
ORF Name	NTID AAID	<u>NT</u> Length	Length	Score	Probability
34094385_c1_23	1864 3784	107	324	155	6.3e-11
Protein name			Locus	Name	Acc#
AttJ			gp:U59	485	1150405 163
Description					U59485:L63 540
Agrobacterium tumes (atrA), AttA1 (attA) AttE (attE), and Atta alternative splice p (attJ), AttK (attK), (attR), AttS (attS),  ORF Name  4770887_f1_3  Protein name hypothetical protein Description  Sulfolobus solfatan	1), AttA2 (att tf (attF) gene products,compl ,AttL (attL),    AttT (attT),    NTID   AAID	cA2), AttB (acceptance), AttB (acceptance), AttB (attM), AttB (attB), MT Length	AttB), AttC cds; AttG cH (attH), AttO (at AA Length 531  Locus gp:SSC	(attC), (attG) g AttI (a tO), Att tV), Att Score [130] Name	AttD (attD), gene, attI), AttJ P (attP), AttR W (attW), AttX  Probability  2.7e-14  Acc# Y18930
Surfolobus soffatal	Ticus 281 kb §			Strain F	
ORF Name	NTID AAID	NT Length	Length	Score	Probability
4875260_c2_33	1866 3786	72	219		
Protein name			Locus	Name	Acc#
Description					
NO-HIT					

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4884702_c1_27	1867	3787	226	681	404	1.4e-37
Protein name NonF					s Name 074603	Acc# AF074603
Description						
Streptomyces grise partial sequence.	us subsp	. griseus	nonactin	biosynt.	hesis ger	necluster,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
6740692_f2_10	1868	3788	165	498		
Protein name				Locus	s Name	Acc#
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
7218752_f3_15	1869	3789	129	390	88	0.030
Protein name					s Name	Acc#
Protein name putative polysaccha	aride po	lymerase			s Name CPS14E	Acc# X85787
<del></del>	aride po	lymerase				
putative polysaccha		lymerase				
putative polysaccha		lymerase AAID	NT Length			
putative polysaccha Description S.pneumoniae cps14	locus.			gp:SP	CPS14E	X85787
putative polysaccha  Description  S.pneumoniae cps14  ORF Name  786305_f1_5  Protein name	NTID	<u>AAID</u> 3790	Length 317	AA Length 954	Score  632  S Name	Probability  9.4e-62  Acc#
putative polysaccha  Description  S.pneumoniae cps14  ORF Name  786305_f1_5  Protein name  probable osmoproted	NTID	<u>AAID</u> 3790	Length 317	gp:SP  AA  Length  954	Score  632  S Name	Probability  9.4e-62
putative polysaccha  Description  S.pneumoniae cps14  ORF Name  786305_f1_5  Protein name	NTID	<u>AAID</u> 3790	Length 317	AA Length 954	Score  632  S Name	Probability  9.4e-62  Acc#
putative polysaccha  Description  S.pneumoniae cps14  ORF Name  786305_f1_5  Protein name  probable osmoproted  Description  ORF Name	NTID 1870 ction bi	AAID 3790 nding pro	Length 317 tein  NT Length	AA Length  954  Locus  pir:G	Score  632  S Name	Probability  9.4e-62  Acc#
putative polysaccha  Description  S.pneumoniae cps14  ORF Name  786305_f1_5  Protein name  probable osmoproted  Description	NTID 1870	AAID 3790 nding pro	Length 317 tein	AA Length  954  Locus pir:G	Score  632 s Name 71892	Probability  9.4e-62  Acc# G71892
putative polysaccha  Description  S.pneumoniae cps14  ORF Name  786305_f1_5  Protein name  probable osmoproted  Description  ORF Name	NTID 1870 ction bi	AAID 3790 nding pro	Length 317 tein  NT Length	AA Length  954  Locus  pir:G  AA Length  447	Score  632 s Name 71892	Probability  9.4e-62  Acc# G71892
putative polysaccha  Description  S.pneumoniae cps14  ORF Name  786305_f1_5  Protein name  probable osmoproted  Description  ORF Name  792090_f2_12	NTID 1870 ction bi	AAID 3790 nding pro	Length 317 tein  NT Length	AA Length  954  Locus  pir:G  AA Length  447	Score  632 S Name 71892 Score	Probability  9.4e-62  Acc# G71892  Probability

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
12273437_f3_58	1872	3792	337	1014 1680	8.3e-173
Protein name				Locus Name sp:SYGA_MORCA	Acc# P77892
Description					
ALPHA CHAIN) (GLYR	S)				<del></del> .
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
14181500_£2_36	1873	3793	693	2082 1593	1.4e-163
Protein name				Locus Name sp:SYGB_HAEIN	Acc# P43822
Description					<del></del>
BETA CHAIN) (GLYRS	)		· · · · · · · · · · · · · · · · · · ·		
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
19650062_c1_101	1874	3794	279	840 581	2.4e-56
Protein name				Locus Name sp:BUDC_KLEPN	<u>Acc#</u> Q48436
Description					
ACETOIN (DIACETYL)	REDUCTA	SE, (ACE	OIN DEHYDR	OGENASE) (AR)	
ORF Name	NTID	<u>AAID</u>	NT Length	AA Score	Probability
21648382_f1_22	1875	3795	279	840 813	6.2e-81
Protein name				Locus Name sp:ACCA_ECOLI	Acc# P30867
Description					·····
(EC 6.4.1.2)					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
21650017_c2_112	1876	3796	254	765 437	4.3e-41
Protein name				Locus Name sp:LPTP_ECOLI	Acc# P23885
Description					
LEUCYL/PHENYLALANY	L-TRNA-	-PROTEIN	TRANSFERAS	Ε,	

ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
21657752_c3_147	1877 3797	345	1038 580	3.0e-56
Protein name			Locus Name sp:YZ37_SYNY3	<u>Acc#</u> Q55480
Description				
HYPOTHETICAL SUGAR	KINASE SLR0537			
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
21987811_£2_34	1878 3798	237	714 384	6.2e-41
Protein name			Locus Name	Acc# P44528
Description				
(EC 2.7.8.5) (PHOS	PHATIDYLGLYCEROPH	OSPHATE S	YNTHASE) (PGP SY	NTHASE)
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
22038132_£3_67	1879 3799	76	231	
Protein name			Locus Name	Acc#
Description				
NO-HIT				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
22384628_f1_5	1880 3800	448	1347 1005	2.8e-101
Protein name			Locus Name sp:YKGC ECOLI	Acc# P77212
Description				
INTERGENIC REGION				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
23493812_f2_33	1881 3801	998	2997 972	6.2e-119
Protein name  metalloprotease 1			Locus Name gp:AF061243	Acc# AF061243
Description		.u. es	<b>-</b>	J
Homo sapiens metal	loprotease 1 (MP1)	mRNA, c	omplete cds.	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
23875027_f2_50	1882	3802	396	1191	621	1.4e-6	0
Protein name					Name UC_HAEIN		Acc# P44433
Description							
(PSEUDOURIDYLATE SY	NTHASE)	(URACIL	HYDROLYAS	E)		-,	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probab	oility
24118802_c3_138	1883	3803	441	1326	1742	2.2e-1	.79
Protein name serine hydroxymethy	Itransf	erase			Name 073769		<u>Acc#</u> AF073769
Description							
Acinetobacter radio complete cds.	resiste	ns serine	hydroxym	ethyltran	nsferase	(glyA)	gene,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probab	oility
24268777_c3_134	1884	3804	1181	3546	1542	1.5e-1	.60
Protein name				Locus	Name		Acc#
Protein name ribonuclease E,:cel protein:message stab protein:RNase E  Description	_		ing	Locus pir:S2		5	Acc# A64852:S45 572:S27311 :A23747:JG
ribonuclease E,:cel protein:message stab protein:RNase E Description	ility-a	AAID	NT Length	Pir:SZ		Probab	A64852:S45 572:S27311 :A23747:JG
ribonuclease E,:cel protein:message stab protein:RNase E Description	ility-a	ltering	NT	pir:SZ	27311	:	A64852:S45 572:S27311 :A23747:JG
ribonuclease E,:cel protein:message stab protein:RNase E Description	NTID	AAID	NT Length	AA Length	27311  Score  145  Name	Probab	A64852:S45 572:S27311 :A23747:JG
ribonuclease E,:cel protein:message stab protein:RNase E  Description  ORF Name  24900257_f1_11  Protein name	NTID	AAID	NT Length	AA Length 261 Locus	27311  Score  145  Name	Probab	A64852:S45 572:S27311 :A23747:JG bility .0
ribonuclease E,:cel protein:message stab protein:RNase E  Description  ORF Name  24900257_f1_11  Protein name  conserved hypotheti	NTID	AAID	NT Length	AA Length 261 Locus	27311  Score  145  Name	Probab	A64852:S45 572:S27311 :A23747:JG Dility .0 Acc# B72287
ribonuclease E,:cel protein:message stab protein:RNase E  Description  ORF Name  24900257_f1_11  Protein name  conserved hypotheti  Description	NTID  [1885]	AAID 3805	NT Length 86	AA Length  261  Locus  pir:B'	Score 145 S Name 72287	Probab	A64852:S45 572:S27311 :A23747:JG Dility .0 Acc# B72287
ribonuclease E,:cel protein:message stab protein:RNase E  Description  ORF Name  24900257_f1_11  Protein name  conserved hypotheti  Description  ORF Name	NTID [1885] cal pro-	AAID 3805 tein AAID	NT Length 86	AA Length Locus pir:B'  AA Length Locus Locus Locus	27311  Score  145  Name 72287	Probab  3.8e-1	A64852:S45 572:S27311 :A23747:JG Dility .0 Acc# B72287
ribonuclease E,:cel protein:message stab protein:RNase E  Description  ORF Name  24900257_f1_11  Protein name  conserved hypotheti  Description  ORF Name  25657776_c1_102	NTID [1885] cal pro-	AAID 3805 tein AAID	NT Length 86	AA Length Locus pir:B'  AA Length Locus Locus Locus	Score  145 S Name 72287  Score  180 S Name	Probab  3.8e-1	A64852:S45 572:S27311 :A23747:JG  pility  Acc# B72287

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26565686_c3_149	1887	3807	348	1047	691	5.2e-68
Protein name hypothetical prot	ein slr0	787		Locus	Name 77001	<u>Acc#</u> S77001
Description	<del></del>			- L		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
26754011_c1_86	1888	3808	357	1074	1765	8.1e-182
Protein name				Locus	s Name	Acc#
NAD repressor/NMN	transpor	rter NadR	p	gp:MC	U73324	U73324
Description						
Moraxella catarrh repressor/NMN tran synthetase alpha s	sporter 1	NadRp (Na	dR) genes,	partial		_
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
2845637_c3_137	1889	3809	172	519	168	1.4e-12
Protein name					Name	Acc#
Description				sp:us	IC_ECOLI	P26602:P76
CHORISMATEPYRUV	ATE LYASI	Ξ,				
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
30332811_£2_51	1890	3810	541	1626	1087	5.7e-110
Protein name				Locus	Name	Acc#
exopolyphosphatas	е			gp:AF	053463	AF053463
<u>Description</u>				<u> </u>		
Pseudomonas aerug complete cds.	inosa th	ioredoxin	(trx) and	l exopolyr	phosphata	se(ppx) genes,
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
30726562_f3_62	1891	3811	768	2307	2256	7.6e-234
Protein name hypothetical prot	ein			Locus	Name	Acc# Y11998
				7 (ab. r. r.	. 02	
Description						
P.fluorescens FC2	.1, FC2.7	2, FC2.3c	, FC2.4 an	a FC2.5c	open rea	dingirames.

ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
33240686_c3_140	1892	3812	260	783 155	1.4e-09
Protein name				Locus Name	Acc#
Description					P31215:P77 227
PNUC PROTEIN		<i>w</i> -			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
34421878_c2_120	1893	3813	408	1227 991	8.5e-100
Protein name				Locus Name sp:YHIN ECOLI	Acc#
Description				Sp. 111111_30033	P37631:P76 705
HYPOTHETICAL 43.8	KD PROT	EIN IN RHS	B-PIT INT	ERGENIC REGION	•
ORF Name	NTID	AAID	<u>NT</u> Length	AA Score	Probability
34578126_f3_54	1894	3814	355	1068 91	0.023
Protein name				Locus Name	Acc#
translation elongat chain PIK-A49:phosph activator PIK-A49			-	pir:A45325	A45325:B45 325:C45325
Description					:D45325:E4
ORF Name	NTID	AAID	NT Length	AA Score	Probability
4695252_f2_29	1895	3815	213	642 497	1.9e-47
Protein name				Locus Name sp:YHGI_ECOLI	Acc# P46847
Description					
HYPOTHETICAL 21.0	KD PROT	EIN IN BIO	H-GNTT IN	TERGENIC REGION	(0191)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> <u>Length</u> <u>Score</u>	Probability
4875885_c2_126	1896	3816	164	495 120	1.7e-07
Protein name				Locus Name sp:YFMU_COXBU	<u>Acc#</u> P45680
Description					
HYPOTHETICAL 15.8	KD PROT	EIN IN FMU	-RPMH INT	ERGENIC REGION	

ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
6651712_f1_15	1897	3817	536	1611	1545	1.7e-	158
Protein name isocitrate lyase				_	s Name 004651		Acc# AB004651
Description							
Hyphomicrobium methors transporter, methion:	_	-		_	-	nicphos	sphate
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
6759625_c3_150	1898	3818	187	564	248	4.6e-	21
Protein name hypothetical protes	in TP0895			Locu pir:D	s Name 71266		<u>Acc#</u> D71266
Description				J [			
ORF Name 14642925_f3_23		AID 3819	NT Length 351	AA Length 1056	Score	Proba	bility
Protein name	ا اــــــا ا			Locu	s Name		Acc#
Description							
NO-HIT		· · · · · · · · · · · · · · · · · · ·					
ORF Name	NTID A	AID	<u>NT</u> Length	<u>AA</u> Length	Score	Proba	bility
16693750_f3_17	1900	3820	100	303	137	2.7e-	09
Protein name conserved hypothet:	ical prote	in yerL		Locus pir:A	s Name 69795		<u>Acc#</u> A69795
Description							
ORF Name 183437_f3_18		AID 3821	<u>NT</u> Length 496	AA Length	Score 2443	Proba	bility 253
Protein name					s Name		Acc# Q49091
Description							
PUTATIVE AMIDASE,					· · · · · · · · · · · · · · · · · · ·	-	

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
1987793_£2_16	1902	3822	264	795 189	8.2e-15
Protein name				Locus Name sp:MINC_ECOLI	Acc# P18196
Description					
CELL DIVISION INHI	BITOR M	INC			
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
22078181_c3_52	1903	3823	137	411 161	7.6e-12
Protein name				Locus Name	Acc#
mai-type protein				pir:D72129	D72129
Description					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
22942053_f1_9	1904	3824	70	210	
Protein name				Locus Name	Acc#
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
23492792_c2_37	1905	3825	203	612 141	1.0e-09
Protein name				Locus Name sp:CYC5_AZOVI	Acc# P11732
Description					
CYTOCHROME C5					
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length Score	Probability
24259651_f2_14	1906	3826	319	960 462	9.7e-44
Protein name				Locus Name	Acc# P32129
Description					
HYPOTHETICAL 36.3	KD PROT	EIN IN DS	BA-POLA IN	TERGENIC REGION	

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
24351556_c1_28	1907	3827	231	696	389	5.3e-36
Protein name Outer membrane p	rotein hom	nolog		_	s Name	Acc# AF067083
Description						<del></del>
Vitreoscilla sp. repressor binding			tial cds;			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
30203430_c2_35	1908	3828	86	261		
Protein name				Locu	s Name	Acc#
<u>Description</u>						
NO-HIT	···				= ····	
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	Score	Probability
31423200_t3_25	1909	3829	181	546	622	1.1e-60
Protein name				Locu	s Name	Acc#
cell division in site-determining		_	n	pir:C	CECID	B31877:D64
Description	-					863
			NTT			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4729837_f3_19	1910	3830	317	954	1626	4.4e-167
Protein name				Locus	s Name	Acc#
BRO-1				gp:MC	BLABRO1	Z54180
Description						
M.catarrhalis bl	a gene.					
ORF Name	NTID	AAID .	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4791053_c1_31	1911	3831	72	219	54	0.0063
Protein name					s Name	Acc#
gag protein				gp:MU	SERVGG2	M26006
Description						
Mouse endogenous 15.3.	retroviru	ıs trunca	ted gag ge	ne, comp.	lete cds,	clonedel env-2

ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length Score	Probability
976677_±3_20	1912 3832	494	1485 2124	7.4e-220
Protein name			Locus Name sp:YBL3_MORCA	<u>Acc#</u> Q49092
Description				
HYPOTHETICAL 46.4	KD PROTEIN IN BLO	R-1 3'REG	ION (ORF3)	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
14469635_£2_5	1913 3833	716	2151 1425	8.7e-146
Protein name			Locus Name	Acc# P44573
Description				
OLIGOPEPTIDASE A,				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
19569430_c3_39	1914 3834	275	828 454	6.8e-43
Protein name			Locus Name	Acc# P75772
Description				
HYPOTHETICAL 28.8	KD PROTEIN IN MOA	E-RHLE IN	TERGENIC REGION	
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
21718878_c1_20	1915 3835	269	810	
Protein name			Locus Name	Acc#
Description				
NO-HIT				
ORF Name	NTID AAID	<u>NT</u> Length	AA Length Score	Probability
22847175_t3_15	1916 3836	83	252 81	0.0023
Protein name			Locus Name	Acc# P56622
Description			<u>-</u>	
HYPOTHETICAL 7.6 K	D PROTEIN IN SLYD	-KEFB INT	ERGENIC REGION	

ORF Name	NTID AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	Score	Probability
23445300_c3_37	1917 3837	923	2772	778	2.2e-100
Protein name			_	Name	Acc#
prolyl oligopeptida	se, precursor		pir:A3	8086	A38086
Description				<u>-</u>	<del></del>
ORF Name	NTID AAID	NT Length	Length	Score	Probability
3907568_c2_28	1918 3838	124	375	70	0.033
Protein name				Name	Acc#
ORF102			gp:AF1	.62221	AF162221
Description					
Xestia c-nigrum gra	nulovirus gen	ome, complet	e sequenc	e.	
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
4773287_c1_26	1919 3839	212	639	503	4.4e-48
Protein name				Name V_ECOLI	Acc# P52061
Description			<u> </u>		
HYPOTHETICAL 21.0 H	D PROTEIN IN	GSHB-ANSB IN	TERGENIC	REGION (	(0197)
ORF Name	NTID AAID	<u>NT</u> Length	<u>AA</u> Length	Score	Probability
964212_c3_35	1920 3840	410	1233	104	0.0091
Protein name				Name	Acc#
voltage-dependent a	nion channel	protein 1b	gp:AF1	.78951	AF178951
Description				-	
Zea mays voltage-de cds; nuclear gene fo			otein 1b (	vdaclb)	mRNA, complete

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